Modification of reactive oxygen species scavenging capacity of chloroplasts through plastid transformation

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Abstract Reactive oxygen species (ROS), including superoxide anions, hydrogen peroxide and hydroxyl radicals are generated through normal biochemical processes, but their production is increased by abiotic stresses. The prospects for enhancing ROS scavenging, and hence stress tolerance, by direct gene expression in a vulnerable cell compartment, the chloroplast, have been explored in tobacco. Several plastid transformants were generated which contained either a Nicotiana mitochondrial superoxide dismutase (MnSOD) or an Escherichia coli glutathione reductase (gor) gene. MnSOD lines had a three-fold increase in MnSOD activity, but interestingly a five to nine-fold increase in total chloroplast SOD activity. Gor transgenic lines had up to 6 times higher GR activity and up to 8 times total glutathione levels compared to wild type tobacco. Photosynthetic capacity of transplastomic plants, as measured by chlorophyll content and variable fluorescence of PSII was equivalent to non-transformed plants.

The response of these transplastomic lines to several applied stresses was examined. In a number of cases improved stress tolerance was observed. Examples include enhanced methyl viologen (Paraquat)-induced oxidative stress tolerance in Mn-superoxidase dismutase over-expressing plants, improved heavy metal tolerance in glutathione reductase expressing lines, and improved tolerance to UV-B radiation in both sets of plants.

Keywords Chloroplast transformation · Superoxide dismutase · Glutathione reductase · Abiotic stress

Introduction

Reactive oxygen species (ROS) are partially reduced, forms of oxygen, routinely produced during normal metabolic processes such as photosynthesis and cellular respiration (Mittler 2002). Light-dependent electron transport processes render the chloroplast a particularly prolific source of ROS, generating superoxide anions (O$_2^•^-$) and hydrogen peroxide (H$_2$O$_2$), which, through transition metal catalysis, can undergo Fenton reactions to liberate the particularly reactive, and highly destructive, hydroxyl radical (OH$^•$) (Lidon and Teixeira 2000). ROS have a high capacity for oxidation of a wide range of cellular components, including lipids, nucleic acids and proteins (Badawi et al. 2004a), disrupting a range of cellular functions and leading to extensive cell damage. During normal metabolism ROS levels are kept in check by a battery of enzymic and non-enzymic anti-oxidants which have evolved for the purpose. The front line enzymes in this process are superoxide dismutases (SODs), which convert O$_2^•^-$ into H$_2$O$_2$. This in turn needs to be removed through the action of catalase (CAT), or through cycles such as the water-water cycle, the
ascorbate-glutathione cycle, or the glutathione peroxidase cycle (Mittler 2002). These cycles employ additional enzymes including ascorbate peroxidase (APX), monodehydroascorbate reductase (MDAR), dehydroascorbate reductase (DHAR), glutathione reductase (GR) and glutathione peroxidase (GPX). Non-enzymatic players include ascorbate (AsA) and glutathione, in reduced (GSH) or oxidised (GSSG) form.

It is well established that a wide range of abiotic stresses, including light, temperature, drought, ionic stresses and UV radiation, disrupt cellular homeostasis leading to substantial increases in ROS levels, such that the scavenging mechanisms may be overwhelmed leading to cell damage (Mittler 2002). Inevitably, therefore, ROS scavenging capacity has been long recognised as a target for genetic engineering (Foyer et al. 1994; Allen 1995; Bhatnagar-Mathur et al. 2008), and over-expression of genes encoding enzymic components of these defences has been explored in a number of studies. The most intensively studied has been the over-expression of SOD. Several cDNAs have been expressed in transgenic plants, and while some gave no apparent improvement in stress tolerance (e.g. Tepperman and Dunsmuir 1990), most did result in enhanced tolerance to one or more of the abiotic stresses examined. Expression of MnSOD (normally located in the mitochondria, but targeted to the chloroplast in most of these studies) resulted in improved frost tolerance in alfalfa (McKersie et al. 1993), cold and oxidative stress tolerance in maize (Van Breusegem et al. 1999), drought (Wang et al. 2005) and salt tolerance in rice (Tanaka et al. 1999), and salt tolerance in poplar (Wang et al. 2010). Levels of the Fe or Cu/Zn SODs, normally associated with chloroplasts, have also been enhanced, resulting in improved cold tolerance in alfalfa (McKersie et al. 1999, 2000), salt and drought tolerance (Badawi et al. 2004b) in tobacco, and methyl viologen (MV—a superoxide-generating herbicide) tolerance in tobacco (Sen Gupta et al. 1993; Slooten et al. 1995) and poplar (Arisi et al. 1998). Other ROS scavenging enzymes have also attracted attention. Catalase over-expression in tobacco resulted in enhanced recovery from high light intensity and MV (Miyagawa et al. 2000), while expression of an ascorbate peroxidase gene resulted in protection against aminotriazole (but not paraquat)-induced oxidative stress (Wang et al. 1999) and heat stress (Shi et al. 2001) in Arabidopsis, and salt and drought stress in tobacco (Badawi et al. 2004a). A similar outcome was achieved through the overexpression of glutathione reductase in tobacco (Aono et al. 1993) and poplar (Foyer et al. 1995). Some investigators have extended the transgenic approach to explore simultaneous over-expression of more than one enzyme associated with ROS scavenging. Thus, Aono et al. (1995) crossed transgenic tobacco plants overexpressing SOD or GR and demonstrated a higher level of tolerance to MV-induced oxidative stress in the hybrids, compared to that of either parent. Similarly, SOD and APX have proven an effective combination in transgenic tobacco (Kwon et al. 2002; Kim et al. 2005), potato (Tang et al. 2006) and tall fescue (Lee et al. 2007a). In one study with tobacco (Lee et al. 2007b) three components of the ROS defence network (SOD, APX and DHAR) were shown to provide further improvements in tolerance to salinity and MV, in comparison with plants over-expressing only the first two of these.

All the above investigations proceeded through nuclear transformation, although most involved targeting of the recombinant enzyme to the chloroplast through the inclusion of a transit sequence. A convenient chloroplast transformation system for tobacco opens the way for the alternative strategy (hitherto unexplored) of examining the interplay of the various components of the ROS defences using direct expression in the chloroplasts. This negates any uncertainty about the efficiency of plastid targeting and, in the absence of any known mechanism for protein export from the chloroplasts, ensures we are exclusively gauging the consequences of over-expression in this organelle. We have embarked on a systematic investigation of the impact of increased activity of ROS scavenging enzymes through direct plastid transformation. The present report describes the results obtained for the first two target enzymes to be explored in this way, MnSOD and GR. SOD was chosen because of its key role in the first step of all the defence pathways, conversion of O2•− into H2O2, and the Mn form selected because, in contrast to the other isoymes, it is not sensitive to inactivation by its reaction product, H2O2 (Bowler et al. 1991). It has also proven an effective choice in previous over-expression studies carried out through nuclear transformation (e.g. McKersie et al. 1993; Tanaka et al. 1999; Van Breusegem et al. 1999). GR was an attractive choice because of its essential role in glutathione homeostasis, specifically in replenishing GSH in both the ascorbate-glutathione and glutathione peroxidase cycles (Mittler 2002).

Materials and methods

Chloroplast transformation vectors

The coding regions of MnSOD and gor were cloned by PCR, using High Fidelity Taq polymerase (Clontech, Saint-Germain-en-Laye, France), and ligated initially into pCR2.1-TOPO (Invitrogen, Paisley, Scotland, UK). PCR products were generated with the addition a NheI and XbaI restriction enzyme site at the 5′ and 3′ ends respectively. PCR products in recombinant clones were fully sequenced, and then the fragments were subcloned into the plastid transformation vector, Nicotiana plumbaginifolia manganese superoxide
dismutase (MnSOD), of 690 bp length, excluding the chloroplast transit peptide sequence, was cloned from the plasmid pHISOD (Bowler et al., 1991, obtained from Professor Dirk Inze, Rijksuniversiteit Gent, Belgium) with the primers 5'-atgctagcgcactacgaaccctagtgagcaga-3' and 5'-gctctagattaacgcca-3'. The Escherichia coli glutathione reductase (gor) sequence (1.4 kbp) was cloned from E. coli genomic DNA with the primers 5'-cggttcgtgaacttcttttc-3' and 5'-gcctgcaagactgttaagtt-3'. The P1 probe (700 bp) was generated with the primers -aagccctggtaaggttcttc-3' and -gctctagattaacgcca-3'.

Fig. 1 Diagram of transgene expression cassettes inserted into tobacco plastid DNA. Map of wild type plastid DNA region of vector insertion and of plasmid pHK-MnSOD and pHK-GR. a Wild type plastid DNA showing the 16SrDNA, trnV, rbcL, and rps12/7 genes along with position of BamHI restriction sites relevant to the Southern blot. Location of the rrm16 probe, P1 is shown. P1 hybridises to a 3.0 kb fragment in the wild type plants, and larger 4.5 and 5.8 kb fragments in the pHK-MnSOD and pHK-GR transformative lines respectively, when total genomic DNA is digested with BamHI. b Location of cloned flanking pieces of tobacco plastid DNA, with the selectable marker gene aadA and the position of the inserted MnSOD or gor downstream of the Prm promoter. MnSOD or gor were cloned into Neilland XbaI sites of pHK34, replacing the neo coding region in each case. Coding regions were immediately downstream of the 3' end of the coding region of the DB of rbcL and driven by the Prm promoter. MnSOD or gor were followed by the rbcL 3' UTR (TrbcL).

Generation of transplastomic plants by particle bombardment

Gene delivery by particle bombardment and the selection process of transplastomic tobacco shoots (Nicotiana tabacum var Petit Havana) clones were performed essentially as described by Daniell (1997). Tobacco leaves were bombarded using a biolistic device (PDS-100/He; Bio-Rad, Hercules, CA). After bombardment, leaves were cut into approximately 25 mm² explants and these were placed on RMOP medium (Svab et al. 1990) containing 500 g ml⁻¹ spectinomycin and green shoots were subcultured onto basal MS (Murashige and Skoog 1962) medium containing 500 g ml⁻¹ spectinomycin. Leaf pieces were placed under at least two rounds of regeneration before regenerated shoots reached homoplasmy. After homoplasmy was confirmed, transplastomic plants were moved to soil. Seed was collected from self-pollinated transgenic plants and used for further analysis.

Southern blot analysis

Southern blot analyses, using 10 µg of BamHI digested genomic DNA, followed by chemiluminescent detection, were carried out as described by McCabe et al. (1997). The P1 probe (700 bp) was generated with the primers 5'-aagccctgtagtctctc-3' and 5'-cggttcctgtaactctttct-3'.

Western blot of MnSOD and GR

Concentration of extracted proteins was determined by a modified Bradford assay (1976) using the BioRad protein assay dye reagent. SDS-PAGE was carried out as described in Sambrook et al. (1989) followed by transfer of proteins to a nitrocellulose membrane (BioRad, Perth, Scotland, UK) (0.2 micron), on a Trans-Blot SD Semi-Dry Electrophoretic Transfer Cell (BioRad Perth, Scotland, UK). MnSOD was detected with a MnSOD polyclonal antibody raised in rabbits against recombinant N. plumbaginifolia MnSOD purified from yeast (Bowler et al. 1991), and kindly provided by Professor Dirk Inze (Rijksuniversiteit Gent, Belgium). GR was detected with a rabbit polyclonal antibody against glutathione reductase, purchased from Lab Frontier (Seoul, Korea) diluted 1:2,000 in blocking buffer. The secondary antibody used was anti-rabbit IgG whole serum peroxidase conjugate (1:2,000 dilution in blocking buffer). Blots were then developed using the chemiluminescent detection system, Super Signal West Pico Kit (Pierce, Aalst, Belgium), according to the manufacturer’s instructions, followed by exposure to Kodak X-ray film.

Leaf material

Leaf material used in all biochemical analyses and stress tests were taken from the 3rd and 4th nodes of 8-week old plants grown in growth rooms at 25°C under a 16 h photoperiod.

SOD analyses

This method was adapted from Chen and Pan (1996). Leaf tissue (100 mg) was frozen in liquid nitrogen and ground to a fine powder in a micro-centrifuge tube. Extraction buffer (0.15 M Tris, pH 7.5, 1% (v/v) Tween 20, 0.5 mM EDTA)
(100 μl) was added and the tissue was ground again and vortexed for 1 min. Samples were then centrifuged at 16,100×g for 15 min at 4°C. The supernatant was transferred to a new microcentrifuge tube and centrifuged again at 16,100×g for 10 min at 4°C to clear any remaining debris. The supernatant was then kept on ice and immediately used for enzyme level determination or stored at −20°C until further use.

Electrophoresis was carried out at 4°C under non-denaturing conditions according to Gabriel (1971) using a 13% separating polyacrylamide gel with a 4% stacking gel in standard tris-glycine buffer (pH 8.3). Samples were electrophoresed at 80 V through the stacking gel for 15 min and 120 V through the separating gel for 60 min. The gel was then soaked in 25 ml of buffer 1 (1.23 mM NBT in potassium phosphate buffer, pH 7.0) for 15 min, briefly washed in dH2O, then soaked in 30 ml buffer 2 (28 mM TEMED, 28 μM riboflavin, in potassium phosphate buffer, pH 7.0), in the dark, with shaking at 75 rpm, for 15 min. Gels were briefly washed in dH2O then illuminated on a light box with a light intensity of 30 µEm−2 s−1 for 15 min. Gels were then photographed using the SynGene gel viewing system. Band identification was according to Bowler et al. (1991).

SOD activity

This method was adapted from Chen and Pan (1996). Leaf tissue (100 mg) was frozen in liquid nitrogen and ground to a fine powder in a microcentrifuge tube. 100 μl extraction buffer (0.15 M Tris, pH 7.5, 1% (v/v) Tween 20, 0.5 mM EDTA) was added and the tissue was ground again and vortexed for 1 min. Samples were then centrifuged at 13,000 rpm for 15 min at 4°C. The supernatant was transferred to a new microcentrifuge tube and centrifuged again at 13,000 rpm for 10 min at 4°C to clear any remaining debris. The supernatant was then kept on ice and immediately used for enzyme level determination or stored at −20°C until further use.

SOD activity was measured using the SOD Assay KitWST from Dojindo Molecular Technologies (MD, USA), following the manufacturer’s instructions (www.dojindo.com/newimages/SODKitTechnicalInformation.pdf), with leaf extracts generated as described above.

Mn SOD specific activity was determined on the basis of cyanide insensitivity (Van Breusegem et al. 1999). Two millimolar KCN was added to the samples in the above assay.

Glutathione reductase assay

Leaf tissue (100 mg) was homogenized in 500 μl extraction buffer (100 mM potassium phosphate (pH 7.5), 1 mM EDTA, 3 mM DDT, 4% (w/v) PVP) in a microcentrifuge tube on ice. The homogenate was filtered through Miracloth (Calbiochem) and centrifuged at 16,100×g for 30 min at 4°C. The supernatant was collected and placed on ice for use immediately in enzyme assays or stored at −80°C. This assay was as described by Foster and Hess (1980), based on the reduction of glutathione (GSSG) by NADPH in the presence of glutathione reductase.

This reaction is measured by the decrease in absorbance at 340 nm using an extinction coefficient (εmM) of 6.22 mM−1 cm−1 for NADPH. One unit of GR activity will cause the oxidation of 1.0 μmol of NADPH at 25°C at pH 7.5. The extracts were used in this assay immediately after extraction. To measure enzyme activity of samples the following were placed into a 1 ml quartz cuvette: 2 mM oxidized glutathione (500 μl), Assay buffer (1 mM EDTA, 100 mM potassium phosphate buffer, pH 7.5) (400 μl), sample extract (50 μl), and 2 mM NADPH (50 μl). The blank consisted of all components listed above except 450 μl Assay buffer was used and no extract was added. The reaction components were mixed by inversion and the absorbance of the sample was read at 340 nm for 2 min at 10 s intervals. The amount of enzyme in the sample was calculated using the following formula:

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\text{GR(units/ml)} = \frac{(\text{rate of change of sample}) - (\text{rate of change of blank})}{\times 6.22 \text{mM}^{-1} \text{cm}^{-1}} \times (\text{volume of sample in ml}).
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Methyl viologen (MV–Paraquat) treatment

Leaf discs (6) of 1.5 cm diameter were cut out of leaves from the 3rd and 4th nodes of soil-grown plants and floated on distilled water. Each disc was then floated individually, abaxial surface down, on 6 ml of water or solutions containing various concentrations of paraquat (0–100 μM) in 35 mm Petri dishes. Leaf discs were then incubated in the dark for 4 h at 23°C. Leaf discs were then placed under a white light source for 48 h with a light intensity of 12μm2/s. Leaf discs were photographed after 24 and 48 h, followed by chlorophyll and electrolyte leakage measurements. Leaf discs (6) were taken from three individual plants of each independent line examined and the assay was repeated three times.

Chlorophyll analysis

Chlorophyll was extracted by soaking leaf discs (1.5 cm in diameter) in 3 ml of 100% methanol for 24 h. The absorbance (A) of the 3 ml solution was then measured spectrophotometrically at 665.2 and 652.4 nm. Total chlorophyll was calculated using the following
equation: Total Chlorophyll (µg/ml plant extract solution) = 1.44A₆₆₅.₂ – 24.93A₆₅₂.₄

Electrolyte leakage measurement

The electrical conductance of the paraquat solutions (C_initial) retained was measured using a conductivity meter (EC215 Conductivity Meter, Hanna Instruments, UK) after the 48 h light treatment. To measure total ion leakage (C_max), leaf discs were autoclaved at 120°C, for 20 min, in their respective solutions and electrical conductance was measured again. Relative electrolyte leakage (EL) was calculated as a percentage: EL = C_initial/C_max * 100.

Ultraviolet (UV-B) radiation exposure

UV treatments consisted of exposure of tobacco leaf discs to UV radiation, generated by Philips TL12 fluorescent tubes (λ_max 315 nm). Leaf discs were exposed to 4.4 W/m² radiation for 16 h, resulting in a total dose of 253 kJ/m². This dose used is unnaturally high (several times that in the southern Europe Mediterranean region) as it was used to probe for potential increases in protection (rather than to do environmental work). The data cannot be directly extrapolated to specific environments as there was no PAR present during the UV-B exposure. The bulb output was filtered through a single layer of cellulose acetate to eliminate UV-C radiation. The irradiance level represents radiation in the spectral range between 295 and 345 nm. Discs were floated on distilled water, with their adaxial side facing the UV source. The decrease in photosynthetic activity was attributed to the UV-B wavelength since the low level of UV-A radiation is ineffective in decreasing chloroplast photosystem II (PSII) activity (Herrmann et al. 1997). Tadić et al. (2001) compared the UV-A and UV-B emitted by TL12 tubes, and showed most emission was in the UV-B range. No additional PAR (photosynthetically active radiation) was applied during the UV treatment. UV levels were measured using a PMA2200 UV-meter (Solar Light Co., Philadelphia, USA). The severity of the stress was assessed by measuring the photosynthetic efficiency of PSII using the saturating pulse fluorescence technique using an imaging PAM (Walz, Effeltrich, Germany). After UV treatments, leaf discs were placed in the dark for at least 30 min before efficiency of PSII was measured. The minimal fluorescence (F0), maximal fluorescence (Fm), and the variable fluorescence (Fv = Fm – F0) were all measured according to van Kooten and Snel (1990). The photochemical yield of open PSII reaction centres, commonly known as the relative variable fluorescence, was calculated as Fv/Fm, as a reflection of the maximal efficiency of PSII. In individual experiments, sampling consisted of 5 discs from three plants per independent line. Overall the experiment was repeated on three occasions.

Effect of cadmium on seed germination

Seed from non-transgenic wild type (WT) and transformed lines were harvested from self pollinated plants grown in soil in growth rooms. Seeds were surface sterilized, and 20 seeds of each line were then placed into sectors on petri dishes containing seed germination medium. Plates were divided into 6 equal sections in the case of GR transformants or 4 sections for MnSOD transformants and seed from all transformed lines from a given gene construct and WT control seeds were placed in rows on a single plate in order to ensure all seeds were treated identically. Triplicate plates were made for each treatment, and three separate experiments were conducted. To test heavy metal tolerance, germination medium containing 0 or 0.5 mM Cadmium(II), added to germination medium as CdCl₂, was used. These plates were wrapped in foil and placed at 23°C for 7 days, after which the foil was removed and plates were then grown in the light for a further 14 days then scored for % germination and photographed.

Statistical analysis

All data are presented as mean (and standard error of the mean). Statistical analysis was performed by one-way analysis of variance (ANOVA). Post-hoc tests used were either Dunn’s or Kruskal–Wallis. Significance level is indicated separately for each experiment.

Results

Transgenic plants expressing MnSOD and gor in chloroplasts

Chloroplast transformants from biolistic transformations were identified by PCR. Three out of 24 spectinomycin resistant shoots regenerated from shoots with pHK-MnSOD were positive for MnSOD by PCR (data not shown), designated as Mn1, Mn2 and Mn6. Similarly, 5 GR transformants, positive for gor integration in the plastid genome, were found from 18 spectinomycin resistant shoots regenerated from pHK34-GR transformations (data not shown), designated GR14, 17, 18, 19 and 20. After two further rounds of tissue culture regeneration on spectinomycin containing medium, genomic DNA was isolated from these 8 shoots and digested with BamHI restriction enzyme. Transplastomic plants were confirmed by Southern blot with the DIG-labelled P1 probe (700 bp) (Fig. 2). Whilst WT tobacco showed a 3.0 kb hybridising band, the
P1 probe hybridised to 4.5 (Fig. 2a) and 5.8 kb (Fig. 2b) bands in the MnSOD and GR transformants respectively. Southern blot analysis confirmed the correct integration of the vector cassette into the plastome and that the plastome of the transformed lines Mn1, 2, and 6 and GR14, 17, 18, 19 and 20 reached the homoplasmic state, in that no WT plastome copies remained after three regeneration cycles on selection medium. WT and transformed plants were transferred to soil and grown to the eight-week stage in growth rooms.

Enzyme analyses

MnSOD and GR protein expression were confirmed by immunoblot analysis of protein extracts separated by SDS-PAGE. The MnSOD antibody detected two polypeptides. Similar amounts of native *N. tabacum* MnSOD were immunologically detected in 50 μg of both WT and transgenic Mn1 soluble leaf extracts as the expected 20 kDa band, while the engineered *N. plumbaginifolia* MnSOD, which was detected in the T1 generation of transgenic line, Mn1 as the expected 22 kDa band (Fig. 3a), due to its fusion to the coding region of the 14 amino acids in pHK34 and not in the wild-type. The engineered *N. plumbaginifolia* MnSOD transgenic band was however, present at a greater intensity than the wild type band representing a significant elevation of the engineered MnSOD protein over the native MnSOD protein.

Expression of the *E. coli* gor was analysed in 20 μg soluble leaf extracts from the T2 generation of each GR transgenic line. The recombinant GR was increased by 2 kDa compared to the native form as a result of the cloning into pHK34. The additional amino acids however did not affect the functional integrity of the protein. Western blots confirmed the presence of the GR protein of the predicted molecular weight, 50 kDa, in leaf extracts of the primary transformants and the T2 generation of each GR transgenic line (Fig. 3b).

The different SOD isoforms were detected by SOD-activity staining of soluble leaf proteins on polyacrylamide gels under non-denaturing conditions (Fig. 3, A, B, C).
A characteristic SOD isoenzyme banding pattern was found for WT tobacco, consisting of two quickly migrating chloroplastic Cu/ZnSOD activity bands, a slower migrating FeSOD and cytoplasmic Cu/ZnSOD, and a much slower migrating MnSOD band (Fig. 3c). In transplastomic lanes a more slowly migrating band was also present corresponding to the engineered MnSOD. Native gel analysis also indicated that cytoplasmic Cu/ZnSOD exhibited similar intensities in both the WT and transplastomic lines, but the latter had greatly increased native MnSOD and chloroplast Cu/ZnSOD. FeSOD also increased, but to a lesser degree, in the three transgenics (Fig. 3c).

SOD enzyme activity was also determined spectrophotometrically. The activity of the endogenous MnSOD in the WT was low, however its activity was higher in transplastomic plants. The engineered MnSOD was present only in transplastomic lines and at equal levels of activity. In primary transformants, line Mn1, showed a ninefold increase in total SOD activity over WT SOD activity and line Mn2 and Mn6, showed a fivefold increase in total SOD activity over WT (data not shown). These increases were confirmed in the T1 generation for each transplastomic line (Fig. 4). Each transplastomic line showed an average threefold increase in MnSOD specific activity. Therefore, differences in total SOD activity of the transplastomic lines were not due solely to differing levels of MnSOD activity but had a contribution from increased activity of the endogenous SODs.

Glutathione reductase and glutathione analyses

GR activity was expressed as nmol NADPH oxidised per min per mg of protein for the T1 progeny of each independently transformed GR line. The GR levels in leaf extracts of plants transformed with gor were substantially higher than the wild type control (Fig. 5a). Transgenic lines GR14 and GR17 had GR levels 6 times higher than WT. GR activity in GR18 and GR20 were 4 and 1.3 times higher respectively, than in the WT. Even the small increase detected in line GR20 was statistically significant ($P < 0.001$). Because GR maintains the levels of cellular glutathione, total glutathione levels were also measured. The increase in GR activity in each transgenic line corresponded to the increase in total glutathione, with GR14 and GR20 increased by 8.3 and 1.8 times respectively compared to WT glutathione levels (Fig. 5b).

Effect of MV treatment on chlorophyll and electrolyte levels

The efficacy of increased expression of MnSOD or GR in tobacco chloroplasts in mediating resistance to oxidative stress was evaluated using MV treatment, with the extent of cellular damage indirectly measured by electrolyte leakage to indicate the extent of membrane rupture. MnSOD transplastomic and WT leaf discs were floated on MV concentrations ranging from 0 to 100 µM. WT leaf discs
were strongly bleached by all levels of MV from 5.0 µM upwards, while all three transplastomic MnSOD lines retained more than 50% of their chlorophyll even at 100 µM MV (Fig. 6a). MV induced approximately 80% electrolyte leakage in WT leaf discs at a concentration of 10 µM, while Mn1 and Mn6 had approximately 20% electrolyte leakage and Mn2 had approximately 30% electrolyte leakage (Fig. 6b). Even at the highest concentration of MV tested, 100 µM, lines Mn1 and Mn2 had lost less than 40% of their electrolytes while Mn6 showed an even greater MV tolerance, having only a 20% loss.

GR lines did not have increased protection from photo-oxidative stress due to MV treatment. Leaves of the same age were sampled and floated on a more limited range of MV concentrations, indicative of the fact that GR transgenic lines as a whole had a greater sensitivity to MV. At a concentration of 5 µM, chlorophyll degradation was visible in the GR lines after 24 h of MV treatment (Fig. 7a), at which point there was little chlorophyll loss in the wild type discs. By 48 h all lines exhibited substantial chlorophyll loss, but the GR lines retained less chlorophyll than WT at 5 µM, indicating greater sensitivity to MV stress (Fig. 7b). When electrolyte leakage was measured, no significant difference in the percent of leakage was seen between GR14, GR20 and WT. However, electrolyte leakage of GR17 and GR18 were significantly greater (*P < 0.001*) than WT (Fig. 7c). The inter-line variation in sensitivity to MV was not correlated with the increase in GR activity or total glutathione levels, as described in Fig. 5.

Differences in UV-B sensitivity in Mn and GR transgenic plants

Leaf discs were exposed to UV-B radiation and its effect on the photosynthetic efficiency of PS II was measured. Leaf discs were exposed to UV-B radiation and initial
experiments were conducted to determine the conditions under which exposure to UV-B induced a significant change in $F_{v}/F_{m}$ in WT and transgenic leaf discs. Leaf discs were exposed to 4.4 W/m$^2$ UV radiations and the $F_{v}/F_{m}$ was measured at 4, 6, 8, 16, and 24 h. An exposure time of 16 h to UV-B was chosen because it was at this time point that significant changes in $F_{v}/F_{m}$ of WT leaf discs were observed. The 16 h exposure results in a total dose of 253 kJ/m$^2$.

Prior to UV-treatment, the $F_{v}/F_{m}$ values were very similar as discussed above. However exposure to 4.4 W/m$^2$ UV-B radiation for 16 h resulted in significant differences in $F_{v}/F_{m}$ values between WT and transgenic lines. WT leaf discs showed a 21% decrease in $F_{v}/F_{m}$ and the degree to which the $F_{v}/F_{m}$ of wild type and Mn transgenic lines were affected by UV-B was extremely significant ($P = 3.4324 \times 10^{-8}$) (Fig. 8a). No statistical difference was observed in the change in $Fv/Fm$ among Mn transgenic lines; exposure of leaf discs of all Mn transgenic lines resulted in a 9% decrease in $F_{v}/F_{m}$ (Fig. 8a). Similar results were observed in the GR transgenic lines in that no significant differences were seen in UV-B tolerance between the GR lines (Fig. 8b), however due to greater variation in individual measurements, results were less significant ($P = <0.001$) than the effect seen in Mn lines. The leaves of all GR plants tested resulted in an 8.2% decrease in $F_{v}/F_{m}$ which is not statistically different from the Mn transgenic lines tested, however it is a substantial improvement over the 21% decrease in WT $F_{v}/F_{m}$ (Fig. 8b).

Effect of cadmium on transgenic tobacco plants

Heavy metal tolerance of T1 generation transgenic seeds was examined by germinating seeds on germination medium containing up to 0.5 mM cadmium (Cd). Only seeds that germinated and continued to grow into seedlings on Cd containing medium, but over 94% of seeds germinated on Cd containing medium were considered fully germinated (Fig. 9a). In the absence of cadmium, WT and transgenic Mn and GR lines showed 100% germination (Fig. 9b), but germination of WT and MnSOD seeds was reduced by at least 80% at 0.5 mM Cd. MnSOD transgenic lines germinated poorly on cadmium containing medium, with an average germination of 14%. However, all five transgenic GR lines showed significantly higher seedling germination percentages over WT and Mn transgenic lines (Fig. 9a, b). Only 23% of wild type seeds germinated on 0.5 mM cadmium containing medium, but over 94% of seeds germinated in lines GR17, GR19, and GR20.

Discussion

This report describes the stable integration and expression of genes encoding two enzymes associated with the scavenging of reactive oxygen species in the chloroplasts of tobacco plants. Immunoblots and an activity gel confirmed the presence and activity of recombinant MnSOD. Total SOD activity was increased sixfold to tenfold in transplastomic lines but much of this was down to an
unexpected elevation of several of the native SOD isoenzymes. Increases in MnSOD specific activity were much smaller; threefold to fourfold in an isoenzyme generally regarded as of relatively low abundance (Van Camp et al. 1997). The bands that are increased in activity correspond to all the organellar SODs (mitochondrial MnSOD, chloroplastic Cu/ZnSOD and FeSOD), while the cytosolic Cu/ZnSOD activity is unaffected. This finding runs counter to some previous observations (e.g. Slooten et al. 1995, McKersie et al. 1999, Kwon et al. 2002), on nuclear transformants, which suggest that over-expression of chloroplast-targeted SODs suppresses the activity of the native enzymes, although this finding is far from universal among the various SOD over-expressing lines. In fact there are two publications which support our observation. Van Breusegem et al. (1999) expressed tobacco MnSOD in maize chloroplasts and published a densitogram of the SOD banding pattern of their plants (Fig. 2c in their paper) which clearly shows elevated activity of all the SOD isoenzyme activities except one, in their transgenic plants. Lee et al. (2007b), reporting on plants simultaneously expressing 2 or 3 ROS scavenging enzymes (including SOD) published an activity gel where there is clear enhancement of an unidentified native SOD (possibly FeSOD on the basis of its proximity to the engineered Cu/Zn enzyme). Curiously the authors of neither paper allude to this interesting finding which, together with our results, suggests redox-mediated retrograde signalling between the chloroplasts and the nucleus. A signalling role of ROS from the chloroplasts, involving induction of nuclear genes encoding defence related proteins, is well established (reviewed in Nott et al. 2006; Pfannschmidt et al. 2009), with H$_2$O$_2$ as the prime candidate. Excess H$_2$O$_2$, produced by stress (or in our case elevated SOD activity) can diffuse from the chloroplast and, probably through mitogen-activated protein kinases (Kovtun et al. 2000), trigger expression of nuclear encoded defence-related genes such as cAPX encoding an ascorbate peroxidise (Kimura et al. 2001), Cat encoding catalase and Gst1 encoding Gluthathione-S-transferase (Polidoros and Scandalios 1999), as well (in the C$_7$-CAM transition species Mesembryanthemum crystallinum) as genes encoding Cu/Zn-, Mn-, and Fe-SOD (Slesak et al. 2003). The current results suggest that increases in H$_2$O$_2$ levels caused by chloroplastic expression of MnSOD, mimic abiotic and biotic stress responses, leading to a similar MAPK induced expression of nuclear-encoded defence proteins, including several organellar SODs, in these transplastomic tobacco lines. Thus the altered levels of oxidative stress tolerance observed are probably not solely the direct result of MnSOD expression in the chloroplasts.

Transplastomic plants expressing MnSOD exhibit substantial increases in tolerance to MV (Paraquat)-induced oxidative stress, as assessed by chlorophyll and electrolyte leakage measurements. This finding is consistent with most of the studies in which plastid SOD levels have been achieved through nuclear transformation and plastid targeting (see Introduction). There are occasional exceptions (e.g. Cu/ZnSOD, Tepperman and Dunsmuir 1990) where no increase in oxidative stress tolerance was achieved in SOD over-expressing plants. Such discrepancies attest to the complexity of the interactions between ROS scavenging enzyme levels (SOD, ascorbate peroxidase, mono- and dehydroascorbate reductase, glutathione-reductase, -peroxidase and -transferase etc), the quantity and redox state of their substrates (ascorbate, glutathione), and the actual

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Fig. 8 Effects of UV-B radiation on the reduction of relative variable fluorescence of wild type and transgenic leaf tissue. Plants were grown in soil in a glass house for 8 weeks. Leaf discs (1 cm in diameter) were excised from leaves of nodes 3 and 4. The minimal fluorescence ($F_0$) and maximal fluorescence ($F_{m}$) were measured on dark adapted samples. Photosynthetic efficiency of PS II was measured as the relative variable fluorescence ($F_v/F_m$), where $F_v = F_m - F_o$. Leaf discs were then exposed to 4.4 Wm$^{-2}$ UV radiation for 16 h. Following UV treatment, $F_v/F_m$ was measured again. The relative variable fluorescence was normalized to that of the non-treated control (WT 100% = 0.692 ± 0.01). a Variable fluorescence of UV-B treated WT and pHK-MnSOD plants. Values represent averages of 40 measurements. Error bars represent standard error of the means. Statistical analysis (Kruskal–Wallis ANOVA) revealed significant differences between Mn1, Mn2, Mn6, and the WT control ($P = 3.4324 \times 10^{-8}$). According to Dunn’s Multiple Comparisons test, no significant differences exist between the three Mn lines. b Variable fluorescence of UV-B treated WT and pHK-GR plants. Statistical analysis (Kruskal–Wallis ANOVA) revealed significant differences between GR14, GR18, GR20 and the WT control ($P < 0.05$).
levels of the individual ROS themselves (some of which are known to have signalling roles), further complicated by the intracellular compartmentation of the enzymes, metabolites and ROS. The consequences of altering an individual component are difficult to predict. For example, Bowler et al. (1991) observed that while large increases in chloroplast MnSOD levels increased oxidative stress tolerance, smaller increases were actually deleterious. Tepperman and Dunsmuir (1990) achieved particularly high expression levels without enhancement of oxidative stress tolerance, but were using a Cu/ZnSOD which, in contrast to MnSOD, is sensitive to the H$_2$O$_2$ generated.

Similar unpredictability is manifest in the oxidative stress response of glutathione reductase (GR) over-expressing transplastomic plants, in which increased GR activity led to increased oxidative stress sensitivity. Elevated GR activity should lead to more reduced glutathione which would generally be regarded as beneficial for redox homeostasis in chloroplasts (Mittler 2002). However, our finding matches that of Creissen et al. (1999) who increased GSH levels by transformation with a gene for a chloroplast-targeted $\gamma$-glutamylcysteine synthetase ($\gamma$-ECS) and found increased oxidative stress, which they explained on the basis of impaired redox sensing in the chloroplast. We contend that the increased MV sensitivity of the transplastomic lines over-expressing GR has a similar basis, although it is not reflected in a study in poplar (Foyer et al. 1995) in which chloroplast targeted GR did result in improved oxidative stress tolerance. Another study with tobacco (Broadbent et al. 1995) showed considerable inter-line variation, some GR over-expressing lines giving a modest increase in oxidative stress tolerance, while others did not. We found that elevation of plastid GR activity over-rides feedback control of glutathione synthesis leading to elevated total glutathione levels. Broadbent et al. (1995) and Foyer et al. (1995) report a similar finding in nuclear transformants of tobacco and poplar respectively.

One previously observed consequence of elevated chloroplast GR, namely tolerance to the heavy metal cadmium (Cd), was also found in the transplastomic GR-overexpressing plants. Cadmium is believed to result in lipid peroxidation (Gallego et al. 1996) as well as disrupting photosynthesis leading to ROS generation (Prasad 1995). Consequently, Pilon-Smits et al. (2000) demonstrated improved Cd tolerance in Brassica juncea plants with elevated GR targeted to the chloroplasts. In the present study a similar clear protective effect was found in germinating transplastomic tobacco seedlings expressing recombinant GR, but not MnSOD, in their chloroplasts. The marked difference between the responses to elevation of these two ROS scavenging enzymes may suggest that the primary role of elevated GR is not in ROS scavenging, but that it is due to the observed increase in the glutathione pool, glutathione being the precursor for the metal-binding peptides, phytochelatins (Cobbett 2000). However, the apparent lack of a protective effect in the MnSOD lines merits further investigation. Cd tolerance was the only stress explored at the seedling level in these lines and there could be major differences in the recombinant protein levels in the chloroplasts of mature leaves and seedlings. Various additional stress tests and analysis of antioxidant levels in these seedlings will be the subject of future studies.

UV irradiation is another stress for which ROS play a major role in mediating damage, and activities of several ROS scavenging enzymes, including both SOD and GR, are boosted by UV-B exposure (Jansen et al. 1998). The observed improvement in UV-B tolerance in response to over-expression of both enzymes in tobacco chloroplasts may again relate to improved ROS scavenging. However, there may be other factors in play. Phytochelatins (synthesised from glutathione) have been shown to have a role

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Fig. 9 Cadmium tolerance test of T1 generation tobacco seedlings overexpressing either glutathione reductase or MnSOD. Seeds of wild type and transgenic lines were germinated on 0.5× MS medium supplemented with 0.5 mM cadmium and grown for 14 days. Approximately 50 seeds of each line were plated out in triplicate in three different experiments. a Representative test of T1 generation GR seeds. b Seed germination and seedling growth after cadmium treatment. Percent of seed that germinated and developed into seedlings on 0.5 mM Cd containing germination medium of total number of wild type and transgenic seeds tested. All standard errors were less than 10% of the mean. GR lines showed increased tolerance to Cd. Mn transgenic lines had no increased tolerance to Cd.
in UV-B tolerance in cyanobacteria (Bhargava et al. 2004), while H$_2$O$_2$ is an important oxidant in phenolpropanoid metabolism, and can therefore influence the synthesis of UV-screening compounds. Manipulation of cytosolic or apoplastic peroxidases has been shown to influence UV-tolerance in tobacco (Jansen et al. 2004; Heggie et al. 2005), so diffusion into the cytosol of excess H$_2$O$_2$, a substrate of peroxidase, generated by MnSOD in the chloroplasts, might also affect the plant’s response to UV. It should be emphasised that this UV protection would be due to alterations in phenolic metabolism in the absence of UV, and is not due to induction of a defence response by UV-B, which would not be plausible during the short time-frame of high level exposure in these experiments.

This report demonstrates that the levels of enzymes associated with ROS scavenging can be modified through direct chloroplast transformation, and that the resulting enhancement of SOD or GR activity results in characteristic alterations in the response to several stresses. We believe that this direct tobacco chloroplast transformation approach has real attractions for further investigations into the roles of the various players in the complex process of oxidative stress amelioration in chloroplasts. The variable results obtained amongst independent nuclear transformants (e.g. Broadbent et al. 1995; McKersie et al. 1999) should be minimised by the targeted gene insertion, via homologous recombination, into the plastome (Svab et al. 1990), eliminating position effects and ensuring uniform expression levels. The lack of any known protein export mechanism from chloroplasts ensures the recombinant protein is localised to the target organelle, and obviates concerns over residual cytosolic expression due to inefficient chloroplast targeting. Furthermore, the importance of multi-component scavenging pathways, such as the ascorbate-glutathione cycle (Mittler 2002), together with the often limited effects of expressing individual components, suggest concerted expression of two or more genes might be more informative, something readily accomplished through chloroplast transformation (Bock 2001; Maliga 2004). This is illustrated by recent experiments in our laboratory (Le Martret et al. 2011) in which we combine expression of dehydroascorbate reductase (DHAR) and glutathione reductase (GR) in tobacco chloroplasts. These two enzymes constitute the parts of the ascorbate-glutathione cycle responsible for regenerating ascorbate, and the co-expressing plants exhibit a pronounced increase in tolerance to methyl viologen-mediated oxidative stress, whereas expression of the individual enzymes has a neutral (DHAR) or negative (GR) outcome. It is anticipated that further investigations of this kind will provide important insights into the interactions between the components of the stress defence armoury of this vulnerable organelle.

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