

Stress-Induced Legume Root Nodule Senescence: Physiological, Biochemical, and Structural Alterations¹

Manuel A. Matamoros, Lisa M. Baird, Pedro R. Escuredo, David A. Dalton, Frank R. Minchin, Iñaki Iturbe-Ormaetxe, Maria C. Rubio, Jose F. Moran, Anthony J. Gordon, and Manuel Becana*

Departamento de Nutrición Vegetal, Estación Experimental de Aula Dei, Consejo Superior de Investigaciones Científicas, Apdo 202, 50080 Zaragoza, Spain (M.A.M., P.R.E., I.I.O, M.C.R., J.F.M., M.B.); Biology Department, University of San Diego, California 92110 (L.M.B.); Biology Department, Reed College, Portland, Oregon 97202 (D.A.D.); and Institute of Grassland and Environmental Research, Plas Gogerddan, Aberystwyth, SY23 3EB, United Kingdom (F.R.M., A.J.G.)

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*Corresponding author; e-mail becana@eead.csic.es; fax 34-976-575620.

Abbreviations: APX, ascorbate peroxidase; DR, dehydroascorbate reductase; γ ECS, γ -glutamylcysteine synthetase; EM, electron microscopy; GR, glutathione reductase; GSHS, glutathione synthetase; hGSH, homoglutathione; hGSHS, homoglutathione synthetase; Lb, leghemoglobin; LM, light microscopy; MR, monodehydroascorbate reductase; N₂ase, nitrogenase; ODB, oxygen diffusion barrier.

Nitrate-fed and dark-stressed beans (*Phaseolus vulgaris*) and peas (*Pisum sativum*) were used to study nodule senescence. In beans, nitrate (1 d) caused a partially reversible decline in nitrogenase (N₂ase) activity and an increase in O₂ diffusion resistance, but minimal changes in carbon metabolites, antioxidants, and other biochemical parameters, indicating that the initial decrease in N₂ase activity is due to O₂ limitation. In peas, dark (1 d) led to a 96% decline in N₂ase activity and sucrose, indicating sugar deprivation as the primary cause of activity loss. In later stages of senescence (4 d with nitrate or 2 to 4 d with dark), nodules showed accumulation of oxidized proteins and general ultrastructural deterioration. The major thiol tripeptides of untreated nodules were homoglutathione (72%) in bean and glutathione (89%) in pea. These predominant thiols declined by approximately 93% after 4 d with nitrate or dark, but the loss of thiol content can be only ascribed in part to limited synthesis by γ -glutamylcysteine, glutathione, and homoglutathione synthetases. Ascorbate peroxidase was immunolocalized primarily in the infected and parenchyma (inner cortex) nodule cells, with large decreases in senescent tissue. Ferritin was hardly detectable in untreated bean nodules but accumulated in the plastids and amyloplasts of uninfected interstitial and parenchyma cells following 2 or 4 d with nitrate, probably as a response to oxidative stress.

Legume N₂ fixation is particularly sensitive to environmental perturbations, including defoliation, water deficit, continuous darkness, and nitrate fertilization (Vance et al., 1979; Witty et al., 1986; Layzell et al., 1990). In most types of stress, the initial decrease of N₂ase activity is associated with a decline in the O₂ concentration reaching the infected cells and bacteroids (Witty et al., 1986; Carroll et al., 1987; Layzell et al., 1990; Escuredo et al., 1996). Prolongation of stress induces premature nodule senescence, which shares some features with natural senescence (nodule aging) such as the loss of N₂ fixation, the increase in lytic activities, and the formation of green pigments from Lb (Pfeiffer et al., 1983; Sarath et al., 1986). This stress-induced senescence has been linked to the enhanced production of oxidants and the lowering of antioxidant defenses (Escuredo et al., 1996; Gogorcena et al., 1997). Oxidants include inorganic (H₂O₂) and organic (lipid) peroxides as well as 'catalytic' iron, which represents the fraction of iron in plant tissues capable of catalyzing the generation of hydroxyl radical through Fenton reactions (Becana et al., 1998).

A major antioxidant mechanism operating in the nodule cytosol is the ascorbate-GSH cycle, which results ultimately in the detoxification of H₂O₂ at the expense of NAD(P)H. The pathway involves the concerted action of four enzymes (APX, DR, MR, and GR) and requires a continuous supply of ascorbate, thiols, and reduced pyridine nucleotides (Dalton et al., 1986; 1992). The initial enzyme of the pathway, APX, may account for up to 1% of the total soluble protein of nodules (Dalton et al., 1998). The thiol tripeptide GSH (γ Glu-Cys-Gly) also participates in the removal of peroxides through the ascorbate-GSH cycle, but it performs additional roles in plants, such as the transport and storage of sulfur, the control of redox status, and the detoxification of heavy metals (Rennenberg, 1995; May et al., 1998). The synthesis of GSH involves two ATP-dependent reactions catalyzed by the enzymes γ ECS and GSHS. Thiol tripeptides are particularly abundant in the leaves, roots, and seeds of legumes, where a thiol tripeptide homolog, hGSH (γ Glu-

Cys- β Ala), may be present in addition to, or instead of, GSH (Klapheck, 1988). Apparently, a specific hGSHS catalyzes the second step of hGSH synthesis in the leaves of some legumes (Macnicol, 1987; Klapheck et al., 1988). It is not known whether hGSH and hGSHS are present in the nodules.

An entirely different antioxidant mechanism, involving the sequestration of catalytic iron by ferritin, may operate also in the nodules. Plant ferritins are composed of 24 subunits and can store up to 4500 atoms of iron in a safe, non-toxic form (Briat and Lobréaux, 1997). In nodules, ferritin may act also as an iron reservoir for N_2 ase, Lb, and other iron-proteins, since ferritin protein increases early in nodulation and then declines concomitantly with the increase in N_2 ase activity and in heme and non-heme iron (Ragland and Theil, 1993). However, little is known about ferritin in senescent nodules. This information may be of considerable interest because nodules are extremely rich in iron (Ragland and Theil, 1993) and this may become available (catalytic iron) for Fenton reactions by proteolysis during nodule senescence, either natural or induced by stress (Becana et al., 1998).

Many other important alterations related to oxygen, carbon, and nitrogen metabolism occur in nodules during senescence, but the precise sequence of these biochemical changes is far from clear (Carroll et al., 1987; Layzell et al., 1990; Gordon et al., 1997). There is also a paucity of information regarding the structural changes involved in stress-induced nodule senescence. LM or EM studies have been carried out with nodules of detopped alfalfa (Vance et al., 1979), dark-treated soybean (Cohen et al., 1986), and nitrate-treated lupine (Lorenzo et al., 1990). Nevertheless, legume symbioses differ in their tolerance to stress and at least some of these differences may be related to the growth pattern (indeterminate versus determinate) of nodules (Sprent, 1980). Comparison of the inhibitory effects of stress on other legumes may provide insight about the mechanisms underlying stress tolerance and nodule senescence. The general objective of this work is to ascertain the time course of events leading to the loss of function and structural deterioration of nodules following stress application to the plant. The specific

objective is to gain further information on the role of some important antioxidants (APX, thiols, and ferritin) in the protection of legume N₂ fixation against the noxious effects of peroxides, free radicals, and catalytic iron. Data presented in this paper are intended to complement two previous reports (Escuredo et al., 1996; Gogorcena et al., 1997), so as to provide an orthogonal comparison of nitrate and dark stress-induced senescence in determinate and indeterminate nodules.

MATERIALS AND METHODS

Plants and Treatments

Nodulated beans (*Phaseolus vulgaris* L. cv Contender x *Rhizobium leguminosarum* biovar *phaseoli* 3622) and peas (*Pisum sativum* L. cv Lincoln x *R. leguminosarum* biovar *viciae* NLV8) were grown in a perlite/vermiculite mixture (2/1) in controlled-environment chambers as described by Gogorcena et al. (1997). When beans had reached the late vegetative stage (30-32 d), pots were divided at random into four groups, which received 10 mM potassium nitrate for 1, 2, or 4 d, whereas control plants receiving N-free nutrient solution were harvested on the third day. Likewise, when peas had reached the late vegetative stage (34-36 d), pots were divided at random into four groups, which were placed in the dark (with otherwise identical conditions) for 1, 2, or 4 d, whereas control plants kept in the light were harvested on the third day. These control plant harvests were arranged so that the maximum age difference between treated and control plants was 2 d. The same protocol was used to produce dark-treated bean and nitrate-treated pea plants so as to obtain nodules for microscopic studies. Nodules to be used for LM and EM analyses were fixed immediately upon detachment as described under "LM Studies". Nodules to be used for biochemical analyses were flash-frozen in liquid N₂ and stored at -80°C for later analysis (within 4-5 weeks).

Nodule Activity and Carbon Metabolites

N₂ase activity and root respiration of intact, undisturbed plants were measured simultaneously using a flow-through gas system (Minchin et al., 1983) housed in a controlled-environment chamber. Root systems were sealed into the growth pots and allowed to stabilize for 18 h. In vivo N₂ase activity was measured as H₂ evolution (Witty and Minchin, 1998) using electrochemical H₂ sensors (City Technology Ltd., Portsmouth, UK) and respiratory CO₂ production was measured using an IR gas analyzer. Measurements were made in air for 5 min and then in a gas stream of 79% Ar/21% O₂. Following exposure to Ar/O₂, steady-state conditions were reached after 60 to 80 min and the external O₂ concentration was then increased over the range of 21 to 60% (8.55 to 24.54 mmol O₂ L⁻¹) in steps of 5 or 10%. Each increase in O₂ took 5 to 6 min and was followed by a 20- to 25-min equilibration period. These data were used to calculate the ODB resistance and carbon costs of N₂ase, as described in Escuredo et al. (1996).

Based on maximum H₂ production under Ar/O₂, the electron allocation coefficient for N₂ was 0.67 for control pea nodules and 0.73 for control bean nodules. The *hup* negative genotype of the two *Rhizobium* strains used in these studies was confirmed by Southern blot analysis of *Eco*RI-digested total DNA using a *hup*-specific DNA probe prepared by dioxigenin-labeling of cosmid pAL618 containing the entire *hup* gene cluster from *R. leguminosarum* biovar *viciae* strain UPM791 (Leyva et al., 1990).

Lb was determined by a method based on the fluorescence emitted by the tetrapyrrol ring after removal of Fe by hot saturated oxalic acid (LaRue and Child, 1979) using myoglobin (horse skeletal muscle; Calbiochem) as the standard. Protein of the nodule cytosol and bacteroids was determined with a commercial dye (Bio-Rad) using crystalline BSA (Sigma) as the standard. Total lipids were extracted from nodules at room temperature essentially as described by Bligh and Dyer (1959).

Carbohydrates were extracted from 0.2 g of nodules with 10 mL of boiling 80% ethanol. The ethanol-soluble extracts were dried in vacuo at 40°C and the soluble compounds redissolved in 4 mL of water. The samples were centrifuged at 20,000g for 10 min and the contents of Glc, Fru, and Suc were determined spectrophotometrically at 340 nm using enzymatic assays coupled to the formation of NADH (González et al., 1995). Starch was extracted from the ethanol-insoluble residue and quantified as the Glc released following digestion with amyloglucosidase (MacRae, 1971).

Catalase and Enzymes of the Ascorbate-GSH Cycle

Antioxidant enzymes were extracted at 4°C from 0.5 g (bean) or 0.25 g (pea) of nodules with a mortar and pestle. Catalase and APX were extracted with 10 mL (bean) or 5 mL (pea) of 50 mM potassium phosphate buffer (pH 7.0) containing 0.5% (w/v) PVP-10. DR, GR, and MR were extracted with 5 mL (bean) or 2.5 mL (pea) of 50 mM potassium phosphate buffer (pH 7.8) containing 1% (w/v) PVP-10, 0.2 mM Na₂EDTA, and 10 mM β-mercaptoethanol. The homogenate was filtered through one layer of Miracloth (Calbiochem) and centrifuged at 15,000g for 20 min.

Catalase activity was assayed by following the decomposition of H₂O₂ at 240 nm (Aebi, 1984). APX and DR activities were determined by measuring the oxidation of ascorbate at 290 nm (Asada, 1984), and the formation of ascorbate at 265 nm (Nakano and Asada, 1981), respectively. MR and GR activities were assayed by monitoring the oxidation of NADH (Dalton et al., 1992) and NADPH (Dalton et al., 1986) at 340 nm, respectively.

All activities were measured at 25°C, in 1-mL reaction mixtures, and within the linear range. Measurements were made with a Lambda-16 spectrophotometer (Perkin-Cetus) during the first 1.5-3 min with no lag period, except for APX, which was measured after a lag of 40 s. Assays were made using sample volumes ranging from 10 μL (catalase) to 100 μL (GR). Where appropriate, controls made by omitting or boiling extracts were run in parallel to correct for nonenzymatic rates,

and buffers and reagents were treated with Chelex resin to avoid contamination by trace amounts of transition metal ions.

Thiols and Thiol Synthetase Activities

GSH and hGSH were extracted from 50 mg of nodules with 500 μ L of 200 mM methanesulfonic acid containing 0.5 mM diethylenetriaminepentaacetic acid. After centrifugation at 13,000g for 5 min, the supernatant was derivatized at pH 8.0 with 2 mM monobromobimane (Fahey and Newton, 1987). The bimane derivatives of GSH and hGSH were separated and quantified by HPLC (Waters) using an analytical Nova-Pak C₁₈ column (3.9 x 150 mm; 4 μ m; Waters) and a 15% methanol/0.25% acetic acid (pH 3.5) solvent at a constant flow of 1 mL min⁻¹. Detection was by fluorescence (model 474 scanning fluorescence detector; Waters) with excitation at 380 nm and emission at 480 nm. Standards of GSH (Sigma) and hGSH (obtained by chemical synthesis at the University of Nebraska, Lincoln) were processed identically to the samples. The proportion of thiol tripeptides present in the oxidized form was determined by an enzymatic recycling procedure using yeast GR to reduce the disulfide forms and vinylpyridine as thiol blocking agent (Griffith, 1980; Law et al., 1983).

The extraction and assays of γ ECS, GSHS, and hGSHS activities were performed by modification of previous protocols, based on the HPLC separation of synthesized γ Glu-Cys, GSH, and hGSH, after derivatization with monobromobimane (Hell and Bergmann, 1988; Kocsy et al., 1996).

Other Metabolites

Ascorbate was extracted from 0.25 g of nodules with 5 mL of 5% (w/v) metaphosphoric acid and quantified by a method based on the ascorbate-dependent reduction of Fe³⁺ to Fe²⁺. Formation of the complex between Fe²⁺ with 2,2'-dipyridyl was measured at 525 nm (Law et al., 1983).

Pyridine nucleotides were extracted from 30 mg of nodules with 2 x 0.5 mL of 0.1 M NaOH (NADH, NADPH) or with 2 x 0.5 mL of 5% TCA (NAD⁺, NADP⁺) at room temperature. After thorough homogenization for 90-120 s in an Eppendorf tube, the extracts were boiled for 6 min, cooled on ice, and centrifuged at 13,000g for 6 min at room temperature. The supernatant (25 μ L for bean, 50 μ L for pea) was made up to 100 μ L with NaOH or TCA, and the nucleotides were quantified by the enzymatic-cycling method of Matsumura and Miyachi (1980).

Oxidant Damage

Lipid peroxides were extracted from 0.5 g of nodules with 5 mL of 5% (w/v) metaphosphoric acid and 100 μ L of 2% (w/v in ethanol) butyl hydroxytoluene (Minotti and Aust, 1987). The extract was filtered through one layer of Miracloth and centrifuged at 15,000g for 20 min. An aliquot of the supernatant was reacted with thiobarbituric acid at low pH and 95°C and cooled to room temperature. The resulting thiobarbituric acid-malondialdehyde adduct was extracted with 1-butanol and quantified by HPLC as described in detail by Iturbe-Ormaetxe et al. (1998).

Protein carbonyl content was measured by derivatization with 2,4-dinitrophenylhydrazine as indicated by Levine et al. (1990) with some modifications. Proteins were extracted from 0.5 g of nodules with 5 mL of 100 mM potassium phosphate (pH 7.0), 0.1% (v/v) Triton X-100, 1 mM Na₂EDTA, and 2.5 μ g each of aprotinin and leupeptin to prevent proteolysis of oxidized proteins during sample preparation. After precipitation of possible contaminating nucleic acids in the samples with 1% (w/v) streptomycin sulfate, an aliquot of 0.8 mL of the extracts was reacted with 0.2 mL of 20 mM dinitrophenylhydrazine (in 2 M HCl) and another aliquot (control) with 0.2 mL of 2 M HCl for 1 h, with vigorous shaking every 10 to 15 min. Proteins were then precipitated with 10% (w/v) TCA and the pellet was washed four times with 1:1 (v/v) ethanol/ethyl acetate. Precipitated proteins were solubilized in 6 M guanidine-HCl (pH 4.5) by incubation for 30 min with shaking. The insoluble material was removed by centrifugation and the absorbance of the hydrazones (derivatized

carbonyls) was measured at 370 nm (Levine et al., 1990). To obtain more accurate results, the amount of protein to be analyzed for carbonyl content was adjusted to 0.5 mg in all samples.

LM Studies

Prior to immunodetection of relevant proteins at the LM or EM levels, bean and pea nodule extracts were subjected to Western analysis (Cresswell et al., 1992). Polyclonal antibodies used in this study were raised in rabbits against N₂ase (universal antibody to the ferroprotein of N₂ase; courtesy of Dr. Paul Ludden), APX (soybean nodule cytosol; Dalton et al., 1993; 1998), and ferritin (soybean seeds; courtesy of Dr. Elizabeth Theil).

Nodules to be used for immunodetection of N₂ase and APX were fixed overnight in 2% paraformaldehyde, 1.25% glutaraldehyde, and 50 mM Pipes (pH 7.2). Fixed samples were dehydrated in an ethanol series, embedded in LR White resin, and cut into 1- μ m sections with a Reichert Ultracut R microtome. For LM immunogold staining, slides were incubated overnight at room temperature in a 1:50 dilution of antibodies raised against N₂ase or APX. For both antigens, the slides were incubated for 1 h with secondary antibody consisting of affinity-purified goat anti-rabbit IgG conjugated to colloidal gold (Auroprobe, Amersham) at a 1:40 dilution. Silver enhancement solution (InsenSEM, Amersham) was applied at room temperature and monitored for sufficient development (15-25 min) following the protocols of the manufacturer. All sections were counterstained with 0.5% safranin in water for 60 s. Representative nodule sections were photographed with a Leitz Laborlux S microscope equipped with a Nikon FE-2 camera.

For LM immunofluorescence staining of APX, the secondary antibody consisted of affinity-purified goat anti-rabbit antibodies conjugated to a Cy3 fluorophore (excitation at 550 nm, emission at 570 nm; Jackson ImmunoResearch Laboratories, West Grove, PA) at a dilution of 1:300 for 30 min. Sections were viewed with a Leitz Laborlux S microscope equipped with a rhodamine filter.

EM Studies

Ultrastructural studies and immunolocalization of APX and ferritin at the EM level were carried out as indicated in Dalton et al. (1993), except that the APX and ferritin antibodies were used at dilutions of 1:500 and 1:25, respectively, and that grids were counterstained only with uranyl acetate. Nodule sections were obtained with a Reichert Ultracut E microtome and representative micrographs for all treatments were taken with a Zeiss 900T transmission electron microscope at 80 kV.

RESULTS

N₂ase Activity and Related Parameters

Beans were treated with 10 mM nitrate for up to 4 d to progressively induce nodule senescence, which was monitored by measuring general markers of metabolic activity (Table I). One day of nitrate application was sufficient to inhibit the *in vivo* N₂ase activity by 73% and to increase the ODB resistance by 3-fold. At this stage, the total root respiration and the carbon cost of N₂ase did not vary, but there were significant increases in Lb and total soluble protein of nodules (Table I). After 2 d with nitrate, the carbon cost of N₂ase increased 1.7-fold whereas Lb and soluble protein returned to control values. After 4 d there was a further increase in the carbon cost of N₂ase up to 2.4-fold and decreases of 53% and 31% in Lb and soluble protein, respectively, relative to the control. An increase in rhizosphere O₂ produced a recovery of N₂ase activity at all stages of the nitrate treatment. Maximum activities at increased O₂ concentrations relative to those at 21% O₂ were 82, 211, 141, and 130% after 0, 1, 2, and 4 d, respectively (Table I).

Dark treatment of peas led to a drastic inhibitory effect on nodule activity. After only 1 d of dark, *in vivo* N₂ase activity was almost abolished, whereas total root

respiration declined by 68% and the ODB resistance was enhanced by 10-fold (Table I). However, there was no effect on the carbon cost of N_2 ase, and Lb and soluble protein only decreased by 29% and 14%, respectively. Prolongation of dark stress for one more day lowered Lb content by an additional 63% but had no effect on total soluble protein. There were no further significant changes after 4 d of darkness, except for total root respiration, which decreased by 86% relative to control (Table I). Increasing rhizosphere O_2 concentrations above 21% caused a slight, but non-significant, rise in N_2 ase activity in peas after 1 d dark, but did not induce any recovery of activity after 2 and 4 d of continuous darkness.

Carbohydrates

Nitrate had considerably less effect on nodule carbohydrates than dark stress. Treatment of bean plants with nitrate led to a progressive decline in Glc, Suc, and starch, whereas Fru increased by 35% after 1 d and declined to 60% of the control value after 2 or 4 d (Table II). The contents of Glc and Suc decreased by 30 to 40% after 1 d of nitrate application, 47 to 57% after 2 d, and 70% after 4 d. Starch was similarly affected after 1 and 2 d with nitrate, but remained at 52% of control after 4 d (Table II).

In contrast, nodules of peas dark-treated for 1 d had lost 97% of their Suc content, along with 69% of Glc, 53% of Fru, and 74% of starch (Table II). The contents of Glc and starch remained essentially constant at this low level following prolongation of the dark treatment to 2 or 4 d, whereas Fru declined by 85% relative to control and Suc virtually disappeared after 4 d (Table II).

Antioxidant Enzymes

Nitrate application for 1 d had only a minor effect on antioxidant enzyme activities from bean nodules, with the exception of GR activity, which increased by 41% (Table III). Following 2 d with nitrate, only APX activity had decreased significantly with respect to control, but after 4 d there was a general decline in

activities of all enzymes of the ascorbate-GSH cycle. The decreases ranged from 18 to 20% for DR and GR to 63% for APX. Catalase activity responded quite differently, with no variation after 2 d of nitrate supply and a 52% decline after 4 d (Table III).

Likewise, after 1 d of dark stress, pea nodules did not show apparent changes in antioxidant activities, but after 2 d there were decreases of 26 to 32% in APX and GR, and catalase activity increased by 31% (Table III). Following 4 d of dark treatment, there were major declines in the activities of the enzymes of the ascorbate-GSH cycle: 40 to 50% for APX, GR and MR, and 82% for DR. In contrast, catalase activity exceeded the control value by 44% (Table III).

Antioxidant Metabolites and Nucleotides

Some improvements of an HPLC method based on the derivatization of thiols to form highly fluorescent adducts (Fahey and Newton, 1987) allowed us to quantify separately, for the first time, GSH and hGSH in nodules (Table IV). Control (untreated) bean nodules contained 103 nmol GSH and 262 nmol hGSH g⁻¹ fresh weight, and pea nodules 698 nmol GSH and 90 nmol hGSH g⁻¹ fresh weight. Hence, GSH was the major thiol (89%) in pea nodules, whereas hGSH predominated (72%) in bean nodules. Assuming uniform distribution and 85% water content in nodules, the total thiol tripeptide (GSH+hGSH) concentrations in bean and pea nodules were estimated as 0.4 mM and 0.9 mM, respectively.

As occurred with antioxidant enzyme activities, nitrate application for 1 or 2 d had only a very limited effect on the content of antioxidant metabolites and pyridine nucleotides in bean nodules (Table IV). After 1 d with nitrate, there were only minor or moderate changes in GSH (37% decrease), ascorbate (20% increase), and NADH and NADP⁺ (18% decrease). The contents of other nucleotides and hGSH remained essentially constant, as did the proportion of the disulfide forms of GSH and hGSH. Following 2 d with nitrate, both GSH and hGSH decreased by 28% but this decrease was not due to simple thiol oxidation since 98% of total thiols were still

present in the reduced form. Only after 4 d was there a significant, although rather modest, increase in the proportion of the disulfide forms, from 2 to 5%. At this stage, GSH and hGSH had declined by 64 and 85%, respectively, and pyridine nucleotides, except NADPH, by 39 to 56%. In contrast, the ascorbate content of nodules treated with nitrate for 4 d was identical to that of untreated nodules (Table IV).

The effect of prolonged darkness on pea nodule antioxidants and nucleotides was also moderate after 1 d, with decreases in the range of 26 to 37% for ascorbate, NAD⁺, and NADH (Table IV). After 2 d, all parameters, except the hGSH content, experienced substantial declines ranging from approximately 24% for NADPH to 62% for NADH. In addition, the proportion of oxidized thiols rose significantly, from 14 to 20%. After 4 d, ascorbate and hGSH declined by 54 to 60%, and GSH by 92%. The nodule content of pyridine nucleotides also decreased markedly, albeit to a lower extent for NAD(H) than for NADP(H). Nevertheless, the NAD⁺/NADH and NADP⁺/NADPH ratios were kept at approximately 3 and 1, respectively, throughout the dark treatment.

Synthesis of GSH and hGSH

Because nitrate and dark treatment led to major declines of GSH and hGSH in nodules which cannot be accounted for by oxidation to the disulfide forms (Table IV), we decided to investigate whether there is GSH and hGSH synthesis in nodules and, in that case, whether this can be impaired by the stress treatments. Optimization of previous HPLC methods enabled us to assay all the enzyme activities required for the synthesis of GSH and hGSH in nodules (Table V), indicating that there is genuine synthesis of thiol tripeptides in the nodule tissue. The first committed step for the synthesis of both tripeptides is catalyzed by γ ECS, and this activity was similar in control (untreated) bean and pea nodules. The second step is thought to be catalyzed by specific synthetases, GSHS or hGSHS.

Both activities were present in bean and pea nodules, but hGSHS activity predominated in bean nodules whereas GSHS activity predominated in pea nodules. This is in agreement with the major thiol tripeptides found in the respective nodules (Table IV), which strongly suggests that the relative abundance of GSH and hGSH is determined, at least in part, by the corresponding thiol tripeptide synthetase activities.

Nitrate supply of bean plants for 2 d led to decreases of 40% in γ ECS activity and 60% in GSHS activity, but had no effect on hGSHS activity; prolongation of treatment up to 4 d led to a decline of 50 to 60% for all three activities of nodules (Table V). Placement of pea plants in the dark for 2 d caused decreases of 40% in GSHS and hGSHS activities of nodules; after 4 d, these activities had decreased by 50 to 60%, whereas the γ ECS activity of nodules had increased by 34% (Table V).

Oxidant Damage

The contents of lipid peroxides (malondialdehyde) and oxidatively modified proteins (carbonyl groups) were used as markers of free radical damage in nodules. The response of both parameters during senescence was, however, distinctly different (Table VI). Nitrate treatment of beans or dark treatment of peas for 4 d caused decreases of 40 to 48% in the content of lipid peroxides of nodules. In contrast, both nitrate and prolonged darkness increased the amount of oxidized proteins in nodules by approximately 34% (Table VI).

Immunolocalization of N₂ase and APX

Immunogold localization of N₂ase with silver enhancement and dark-field LM indicated the presence of abundant protein in the infected cells of control (untreated) bean (Fig. 1A) and pea (data not shown) nodules. Little-or-no labeling occurred in nodules that had been exposed to either of the stress treatments for 4 d (Fig. 1B). Using the same technique but with bright-field LM, APX protein was found to be localized predominantly in the endodermis and adjacent cell layers of

the nodule parenchyma (inner cortex) as well as in the infected zone of control bean nodules (Fig. 2A). Using immunofluorescence with the secondary antibody conjugated to the fluorophore Cy3, a similar distribution of APX protein was noticed in control pea nodules (Fig. 2C). Treatment with either nitrate or dark led to a substantial decrease in labeling intensity in both bean and pea nodules (Fig. 2, B and D), in agreement with the observed declines in enzyme activity (Table III). As expected, bean and pea nodule sections in which rabbit normal serum was used in place of the primary antibody showed only a very sparse background labeling.

The pattern of immunolocalization of APX at the EM level was similar to that observed at the LM level. No label was evident in negative controls (Fig. 3, A and B), whereas strong labeling was present in the cytosol of the parenchyma and infected cells of control bean and pea nodules (Fig. 3, C and D). Label was also noted in the cytosol of interstitial cells (Fig. 4B), over the symbiosomes, and occasionally over the mitochondria and bacteroids (Fig. 4C). Immunolabeling of APX decreased in bean and pea nodule tissue with nitrate and dark stress. The location of labeling was not affected by any of the two treatments.

Ultrastructural Studies

Ultrathin sections of representative nodules from nitrate-treated beans and dark-treated peas were also examined by EM to follow the structural changes occurring during senescence and complement the physiological and biochemical data. In addition, nodules from dark-treated beans and nitrate-treated peas were processed in parallel to allow for orthogonal comparisons.

Infected cells of control (untreated) bean nodules were densely packed with symbiosomes, each enclosing one to four bacteroids. Bacteroids contained abundant poly- β -hydroxybutyrate granules, and organelles of the infected cell, including mitochondria and plastids, were confined to the periphery of the cell. Bean nodules treated for 1 or 2 d with nitrate essentially showed features similar to control nodules (Fig. 4D). After 4 d of nitrate treatment, significant disruption of the

host cytoplasm and of symbiosome membranes was observed; however, there were no evident changes in bacteroids or in poly- β -hydroxybutyrate content (Fig. 4E).

Dark stress resulted in observable changes in bean nodules after only 1 d of treatment (Fig. 4F). Many infected cells, particularly toward the interior of the nodule, showed disrupted cytoplasm and discontinuities in the symbiosome membrane. After 2 d of dark, breaks in the symbiosome membrane were more frequently observed, and cytoplasmic breakdown of infected cells was evident (Fig. 4G). Cell disruption was observed throughout the infected zone of the nodule after 4 d of dark treatment and no intact symbiosomes were evident; however, the integrity of bacteroids was not affected and poly- β -hydroxybutyrate granules were comparable in appearance to those observed in controls (Fig. 4H).

Control pea nodules contained abundant symbiosomes within the dense host cytoplasm. No poly- β -hydroxybutyrate was observed within the bacteroids. Cell organelles were located at the periphery of infected cells, and profiles of the rough endoplasmic reticulum and Golgi apparatus were often apparent. One day of dark resulted in minimal change in infected cells (Fig. 5C). Following 2 d of dark treatment, cytoplasmic disruption was widespread, including cytoplasmic breakdown and lesions in the symbiosome membranes (Fig. 5D). Bacteroids ranged in appearance from those showing slight cytoplasmic disruption to others showing extensive breakdown. After 4 d of dark, bacteroids were often misshapen with disrupted cytoplasm and symbiosome membranes (Fig. 5E). The host cell cytoplasm was not discernible and host cell organelles were rarely observed.

One day of nitrate treatment had little effect on the appearance of infected cells in pea nodules. Cell plasmolysis and discontinuities in the symbiosome membrane were apparent after 2 d with nitrate (Fig. 5F). Bacteroids with irregular membranes were occasionally observed (Fig. 5G). Cytoplasmic and bacteroid degeneration was extensive following 4 d of nitrate treatment, and cellular detail was difficult to discern (Fig. 5H).

Ferritin Localization

Preliminary immunoblots of ferritin showed that the antibody raised to soybean seed ferritin recognized a single protein band of 28 kD, characteristic of ferritin subunits, in bean nodule extracts. As expected, a positive control, consisting of a soluble extract of tobacco leaves overexpressing soybean ferritin, produced an identical band (Fig. 6). The immunoblots also revealed that ferritin accumulated in bean nodules in response to nitrate treatment (Fig. 6).

The soybean antibody, however, exhibited very poor reactivity with pea ferritin (data not shown). Consequently, ferritin was only immunolocalized in bean nodules. Control (untreated) bean nodule sections showed little labeling for ferritin (Fig. 7A) but this was clearly visible after 2 or 4 d of nitrate treatment (Fig. 7, B-D), confirming blot analysis. After 2 d with nitrate, ferritin was localized in the plastids and amyloplasts of uninfected and infected cells, and occasionally over the bacteroids. The heaviest labeling was observed in the amyloplasts of the uninfected interstitial cells and of the parenchyma cells (Fig. 7, B and C). In some sections, scattered arrays or small clusters of ferritin particles could be seen in the amyloplasts without the assistance of gold labeling (Fig. 7B). After 2 d and, especially, 4 d of nitrate treatment, large ferritin aggregates were easily observed in the amyloplasts (Fig. 7, C and D). Quite often, these large deposits did not show any immunolabeling, although it was observed in plastids or amyloplasts within the same nodule sections (Fig. 7C), indicating that the antibody was recognizing isolated ferritin particles and smaller ferritin deposits.

DISCUSSION

In this work we have obtained simultaneously an array of physiological, biochemical, and structural data of bean and pea nodules induced to senesce by treating the plants with nitrate or prolonged darkness. This allowed us to monitor the progress of nodule senescence in a controlled manner and to discern more readily between the reversible and irreversible stages of stress application.

Nitrate-Induced Legume Nodule Senescence

Nitrate had a two-stage effect on bean nodules. In the first stage (1 to 2 d), there were major declines in the *in vivo* N₂ase activity and increases in the ODB resistance and carbon cost of N₂ase, but only moderate or no changes in the nodule content of carbohydrates, antioxidants, and pyridine nucleotides; furthermore, there were significant increases in Lb, soluble protein, and ascorbate. At this early stage, there were no detectable changes in the nodules at the ultrastructural level. Therefore, it would appear that the decrease of N₂ase activity after 1 to 2 d may be attributed to O₂ limitation at the ODB level and not to biochemical factors such as degradation of N₂ase or Lb, oxidative damage of cell components, or sugar deprivation of host cells or bacteroids. However, the causal relationship between N₂ase activity and ODB operation cannot be fully examined by the present data and it is possible that N₂ase activity is being decreased by an, as yet, unknown mechanism that, in turn, results in closure of the ODB. The O₂ limitation through the ODB was confirmed by the partial recovery of N₂ase activity in nitrate-treated plants upon increasing the concentration of rhizosphere O₂. This recovery, and the absence of structural and biochemical damage, indicate that at this early stage the effect of nitrate is still reversible. In contrast, in the second stage (4 d), there was almost completely loss of both N₂ase activity and protein, along with a marked decline in Lb, soluble protein, sugars, antioxidants (except ascorbate), and nucleotides. There was also a general deterioration of nodule

ultrastructure, in particular of symbiosome membranes, and accumulation of oxidized proteins. This stage of nitrate inhibition would appear therefore to be essentially irreversible. However, these conclusions await final verification through recovery experiments.

The effects of nitrate on bean and pea nodules can be compared with the limited information available for other legumes. In lupine and clusterbean (*Cyamopsis*) nodules, N_2 ase activity (assayed in both cases with a closed system) was inhibited by 35% after 3 to 5 d with 20 mM nitrate, and structural degradation of lupine nodules was only recognizable after 10 d (Lorenzo et al., 1990; Swaraj et al., 1993). In bean and pea nodules, N_2 ase activity (assayed in both cases with a flow-through gas system) was inhibited by approximately 85% after only 2 d with 10 mM nitrate. Structural data revealed, in turn, differences in the progression of senescence between pea and bean nodules. In pea nodules, the plasmolysis of host cells and the disruption of symbiosome membranes and bacteroids were already evident after 2 d, whereas, in bean nodules, nitrate had little effect on the shape and poly- β -hydroxybutyrate content of bacteroids even after 4 d. These data suggest that pea nodules are particularly sensitive to nitrate.

Dark-Induced Legume Nodule Senescence

Prolonged darkness severely affected pea nodule metabolism. After only 1 d dark there were major effects on total root respiration, ODB resistance, and carbon costs of N_2 ase. In addition there were moderate decreases in Lb and in some antioxidants and nucleotides; however, the most affected parameters were, by far, the *in vivo* N_2 ase activity and the nodule Suc content, which decreased by 97%. The almost complete depletion of Suc, the major carbon and energy source for host cell metabolism, along with the substantial decreases in other carbohydrates, is the most likely cause for the limitation of N_2 ase activity in dark-treated peas. This conclusion is supported by immunoblots of N_2 ase showing no loss in the protein after 1 d dark in both bean (Gogorcena et al., 1997) and pea (data not shown), and

is also consistent with the finding that isolated bacteroids from dark-treated soybean retained 50% of the initial N_2 ase activity (Sarath et al., 1986) and fully recovered it upon addition of succinate (Carroll et al., 1987). After 2 d dark, pea nodules showed drastic decreases in Lb and carbohydrates, moderate decreases of many antioxidants, accumulation of modified proteins, and evident symptoms of structural deterioration. After 4 d dark, there was a general collapse of metabolism and extensive structural damage.

There are some significant differences in the response of other legume nodules to dark stress. Thus, inhibition of N_2 ase activity was much less dramatic in soybean (Sarath et al., 1986) and clusterbean (Swaraj et al., 1994) than it was in bean (Gogorcena et al., 1997) and pea (this work). Also, 2 d of dark had no significant effect on Lb or total soluble protein in nodules of soybean (Pfeiffer et al., 1983; Gordon et al., 1993), clusterbean (Swaraj et al., 1994), and bean (Gogorcena et al., 1997). However, in pea nodules, the same treatment caused a 74% decline in Lb but only a 14% decline in total soluble protein, which is indicative of a relatively high sensitivity of pea Lb to degradation compared with other cytosolic proteins. This conclusion agrees with an early observation that exposure of peas to 3 d of dark was sufficient to induce greening of nodules and breakdown of 50% of total heme (Roiponen, 1970). Because the pathway for Lb degradation in vivo is largely unknown, it can only be speculated that the rapid loss of pea Lb in dark-stressed nodules is due to a particularly rapid activation or decompartmentation of proteases, which are located in the infected cells and display a high affinity for Lb, especially at the acidic intracellular pH of senescing nodules (Pladys et al., 1991).

Thiols, APX, and Ferritin in Senescent Legume Nodules

Three important antioxidants, GSH and APX, which are critical for the operation of the ascorbate-GSH cycle, and ferritin, which is critical for the control of the cellular concentration of catalytic iron, have been studied in further detail to extend earlier studies on the mechanism of stress-induced nodule senescence.

An enzymatic method (Griffith, 1980) was previously employed to estimate the concentration of GSH in nodules. Because this method could not distinguish between GSH and hGSH (Klapheck, 1988) and it was uncertain whether hGSH was present in nodules, we estimated the concentration of total tripeptides in pea and bean nodules as 0.9 mM (Escuredo et al., 1996; Gogorcena et al., 1997). Using HPLC, we have found in this work that hGSH is present in nodules at variable concentrations: 0.12 mM GSH and 0.31 mM hGSH for bean nodules, and 0.82 mM GSH and 0.11 mM hGSH for pea nodules. Thus, the enzymatic method tends to overestimate the thiol content in plant tissues having primarily hGSH because the reaction of yeast GR with hGSH is faster than with GSH (Klapheck, 1988). For pea nodules, which mainly contain GSH, both methods yielded similar results.

An obvious question raised in this work is why hGSH is the major thiol in bean nodules. Although this cannot be answered at present, our results show that the relative abundance of GSH and hGSH in nodules is likely to be dictated by the presence of specific tripeptide synthetases: hGSHS in bean nodules and GSHS in pea nodules. This conclusion is reinforced by the partial purification of distinct enzymes from pea (GSHS) and mungbean (hGSHS) leaves (Macnicol, 1987), which contain only GSH and hGSH, respectively (Klapheck, 1988). The fact that hGSH is the only thiol tripeptide present in the leaves of some legumes also implies that the corresponding chloroplasts have a functional ascorbate-hGSH pathway to avoid photooxidative damage and that hGSH and GSH share at least an antioxidative role *in vivo*.

The drastic decreases of the major thiols after 4 d treatment, 85% hGSH in bean nodules and 92% GSH in pea nodules, cannot be accounted for by oxidation to the disulfide forms or by the 50 to 60% decline in GSHS and hGSHS activities. Nor can it be ascribed to a limitation of γ ECS activity, either directly (the activity only declined by 60% in bean nodules and even increased moderately in pea nodules) or through the availability of Cys (the content of this substrate decreased by 50 to 60% in both bean and pea nodules after 4 d treatment). The lack of evidence for the

inhibition of thiol synthesis, together with the extensive Lb degradation and oxidative reactions taking place in stressed nodules, which are manifest by loss of Lb heme and accumulation of oxidized proteins, may provide a clue for the large decline in GSH and hGSH. These thiols may be consumed by nonenzymatic reactions with activated oxygen species or by enzymatic degradation, with possible formation of mixed disulfides between thiols and proteins (Rennenberg, 1995). These aspects of thiol catabolism in nodules, and in plants in general, remain virtually unexplored.

Another antioxidant, APX, is critical for the disposal of H_2O_2 in nodules. The APX protein was predominantly located to the parenchyma and infected zone, confirming a previous report in which APX protein and mRNA were shown to be enhanced in the parenchyma and infected cells of alfalfa nodules (Dalton et al., 1998). In the same study, APX protein was found also to be increased in the nodule parenchyma and infected zone of several determinate nodules (Dalton et al., 1998). Our EM studies corroborate the heterogeneous distribution of APX within nodules and indicate that the protein is very abundant in the cytosol of infected cells. The observation of occasional labeling of APX in the mitochondria would explain the detection of enzyme activity in purified mitochondria from soybean nodules (Dalton et al., 1993). The activity and content of APX protein largely decreased in senescent nodules. This may cause, in turn, a lowering in protection against H_2O_2 generated by the respiratory activity of the nodule parenchyma and infected cells.

Finally, we have conducted studies to localize ferritin in bean nodules and to determine the changes in ferritin content during senescence. Ferritin was hardly detectable in control nodules but accumulated in nodules treated with nitrate for 2 or 4 d. The very low content of ferritin protein found in mature, untreated bean nodules is in agreement with the observation that ferritin accumulates in young soybean nodules but declines in mature nodules, when iron storage is apparently no longer required (Bergersen, 1963; Ragland and Theil, 1993). Ferritin was predominantly found in the plastids and amyloplasts of uninfected cells in the infected zone and in the parenchyma cells. The subcellular location of ferritin in bean nodules is fully

consistent with other reports showing ferritin particles in plastids and amyloplasts of soybean, alfalfa, and lupine nodules (Bergersen, 1963; Lucas et al., 1998), in amyloplasts of soybean cell cultures (Briat and Lobréaux, 1997), and in chloroplasts and other plastids of several plants (Seckback, 1982).

The scattered arrays and large deposits of ferritin particles observed after 2 or 4 d with nitrate closely resemble, respectively, the F-2 and F-3 types described by Seckback (1982). The latter category is defined as paracrystalline ferritin arrangements (sometimes including small zones of crystalline structure), such as those observed in plastids of iron-treated *Xanthium* without the assistance of gold labeling. Our finding that the paracrystalline deposits of ferritin-like material did not label as densely as would be expected with the soybean antibody requires further investigation, but similar ferritin structures, clearly immunolabeled, accumulate in the cortex of senescing soybean and lupine nodules (Lucas et al., 1998).

Ferritin synthesis in plants is regulated by iron and is induced by various adverse conditions, including iron overload, which lead to oxidative stress (Briat and Lobréaux, 1997). Recent experiments with de-rooted maize plantlets have shown that H₂O₂ induces ferritin mRNA accumulation in the presence of low iron concentrations and that this effect is prevented by pretreatment of plantlets with antioxidants, indicating that the induction of ferritin gene expression in this system requires an oxidative step (Briat and Lobréaux, 1997). Based on these observations, our results showing ferritin protein accumulation in senescent bean nodules may be interpreted as a response to oxidative stress. This oxidative stress is evidenced by the accumulation of damaged proteins in nitrate-treated bean nodules and is probably a consequence of the lowering of antioxidant activities and the release of catalytic iron from proteins (Becana et al., 1998). Further work is needed to establish the mechanism for ferritin induction in nitrate-treated bean nodules and to determine whether a similar phenomenon occurs in other legume nodules under different types of stress.

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Legends of Figures:

Figure 1. Immunolocalization of N₂ase in bean nodules using dark-field LM. A, Control (untreated) nodules. Bright spots in the infected zone of this section indicate the presence of N₂ase. B, Nodule after 4 d with nitrate. Note the almost complete lack of staining in this section. NC, nodule cortex; NP, nodule parenchyma; INF, infected zone. Bars= 100 μm.

Figure 2. Immunolocalization of APX in nodules of bean (A and B, bright field immunogold detection) and pea (C and D, immunofluorescent Cy3 detection). A and C, Control (untreated) nodules showing strong labeling in the infected zone and a prominent band in the nodule parenchyma. B, Nodules after 4 d with nitrate. D, Nodules after 4 d of dark stress. B and D show a marked decrease in labeling in both the infected zone and nodule parenchyma. V, vascular bundle. Other symbols are as in Figure 2. Bars= 100 μm.

Figure 3. Immunogold localization of APX in control (untreated) nodules. A and B, Negative controls (normal rabbit antiserum in place of APX antibody) of bean and pea nodules, respectively, showing no label in the cytosol or organelles of infected cells. C and D, Bean and pea nodule sections, respectively, showing APX label over the cytosol of infected cells. *b*, bacteroid; *w*, cell wall; *m*, mitochondrion; *p*, plastid. The bar indicated in panel A equals 0.2 μm for all panels.

Figure 4. Ultrastructural changes and APX localization in bean nodules after nitrate and dark treatments. A, Negative control showing no label in the cytosol, organelles, or symbiosome of a control (untreated) nodule. B and C, Control nodules showing APX localization in uninfected and infected cells. Label is seen frequently in the cytosol of both uninfected (B, arrowhead) and infected (C, arrowhead) cells, over

the symbiosome and, occasionally, over mitochondria (C, double arrowhead) and bacteroids (C, asterisk). D, Detail of infected cell of nodule after 2 d of nitrate treatment. Cells and bacteroids are similar in appearance to controls. E, Detail of infected cell of nodule after 4 d with nitrate showing the absence of symbiosome membrane, but little change in bacteroids or in poly- β -hydroxybutyrate granules. F, Detail of infected cells of nodule after 1 d of dark stress showing disruption of the cytoplasm (arrowhead). Symbiosome membranes are disrupted in both of the cells shown. G, Infected cells of nodule after 2 d of dark stress showing lesions in the symbiosome membrane and the cytoplasmic disruption observed throughout the nodule. H, Nodule following 4 d of dark stress showing general disruption of infected and uninfected cells. However, most bacteroids remain intact. *m*, mitochondrion; *p*, plastid; *px*, peroxisome; and *u*, uninfected cell. Bars= 0.4 μ m.

Figure 5. Ultrastructural changes and APX localization in pea nodules after nitrate and dark treatments. A, Negative control showing no label in the cytosol, wall, or symbiosome of an untreated nodule. B, Infected cell of untreated nodule showing distribution of APX label over the cytosol and symbiosome. Label was also observed in uninfected cells and, occasionally, over mitochondria. C, Infected cell of nodule after 1 d of dark stress showing only minor disruption of symbiosome membrane (arrowhead). D, Low magnification micrograph of cell after 2 d of dark stress showing the extent of cytoplasmic disruption and bacteroids with various degrees of breakdown. E, Infected cell of nodule after 4 d of dark stress showing disruption of symbiosome membranes and misshapen, disrupted bacteroids. F-G, Infected cells of nodules after 2 d with nitrate showing some cell plasmolysis (F, arrowhead), discontinuity in the symbiosome membrane, and misshapen bacteroids (G, arrowhead). H, Infected cell of nodule after 4 d with nitrate showing the extensive cytoplasmic and bacteroid degeneration characteristic of this stage. A, B, C, E, F and H, bars= 0.4 μ m. D and G, bars= 2 μ m.

Figure 6. Western analysis of ferritin in bean nodules using chemiluminescence detection. Lanes 0, 1, 2, and 4 were loaded with soluble extracts (20 μg protein) from nodules treated with nitrate for 0, 1, 2 and 4 d. Two extracts obtained independently were loaded for each treatment. Lane T was loaded with a soluble extract (20 μg protein) of transgenic tobacco leaves overexpressing soybean ferritin (Van Wuytswinkel et al., 1998), which served as a positive control. In all lanes a single band at 28 kD, characteristic of ferritin subunits, was observed.

Figure 7. Electron micrographs of bean nodules showing ferritin localization. A, Control (untreated) nodule section showing scant deposition of gold label over infected and uninfected cells. B, Detail of parenchyma cell after 2 d with nitrate showing ferritin localization in plastids and amyloplasts. A linear array of ferritin particles (arrowhead) is visible in the amyloplast. C, Detail of parenchyma cell after 2 d with nitrate showing large ferritin deposits not labeled with gold (arrowhead) as well as disperse labeling of ferritin particles (double arrowhead) in the amyloplast. D, Amyloplast from parenchyma cell after 4 d with nitrate. A large aggregate of ferritin (arrowhead) is present, but did not label with gold. *b*, bacteroid; *p*, plastid; *ps*, peribacteroid space; *s*, starch grain; *v*, vacuole; *w*, cell wall. Bars= 0.4 μm .

Table I. *Some general markers of metabolic activity in senescing legume nodules*

Parameter ^a	Bean nodules (days of nitrate)				Pea nodules (days of darkness)			
	0	1	2	4	0	1	2	4
N ₂ ase activity ^b	1.66 a	0.45 b	0.29 b	0.20 b	0.76 a	0.03 b	0.00 b	0.00 b
N ₂ ase activity in O ₂ ^c	1.36 a	0.95 b	0.43 c	0.26 c	0.56 a	0.04 b	0.00 b	0.00 b
Total root respiration	5.24 a	4.76 a	5.33 a	5.59 a	3.23 a	1.03 b	0.63 bc	0.46 c
ODB resistance	0.67 a	1.98 b	1.87 b	1.90 b	1.47 a	15.01 b	ND ^d	ND
Carbon cost of N ₂ ase	1.90 a	2.10 ab	3.30 bc	4.56 c	2.02 a	2.36 a	ND	ND
Lb	221 a	260 b	205 a	102 c	121 a	86 b	32 c	22 c
Soluble protein	12.4 a	16.0 b	12.1 a	8.6 c	14.2 a	12.2 b	12.2 b	12.9 ab

^aMeans ($n= 4-8$) were compared by one-way analysis of variance and the Duncan's range test. For each parameter and legume species, means denoted by the same letter do not differ significantly at $P<0.05$. Units: N₂ase ($\mu\text{mol H}_2 \text{ min}^{-1} \text{ plant}^{-1}$); respiration ($\mu\text{mol CO}_2 \text{ min}^{-1} \text{ plant}^{-1}$); carbon cost ($\text{mol CO}_2 \text{ mol}^{-1} \text{ H}_2$); ODB resistance ($\text{s cm}^{-1} \times 10^{-6}$); Lb ($\text{nmol g}^{-1} \text{ fresh wt}$); and protein ($\text{mg g}^{-1} \text{ fresh wt}$). ^bMaximum H₂ production under Ar/21% O₂. ^cMaximum H₂ production at increased O₂ concentrations following the Ar-induced decline. ^dND=not determined.

Table II. Carbohydrate content in senescing legume nodules

Metabolite ^a	Bean nodules (days of nitrate)				Pea nodules (days of darkness)			
	0	1	2	4	0	1	2	4
	<i>μg g⁻¹ fresh wt</i>							
Glc 173 a	104 b	92 b	51 c	348 a	109 b	125 b	94 b	
Fru 89 a	120 b	53 c	52 c	70 a	33 b	11 c	10 c	
Suc4445 a	3087 b	1902 c	1111 d	8946 a	313 b	207 b	34 c	
Starch	1095 a	652 b	466 b	521 b	8002 a	2084 b	1526 b	1235 b

^aStatistical analysis of means ($n=3$) was performed as for Table I.

Table III. Antioxidant enzyme activities in senescing legume nodules

Enzyme ^a	Bean nodules (days of nitrate)				Pea nodules (days of darkness)			
	0	1	2	4	0	1	2	4
	<i>μmol min⁻¹ g⁻¹ fresh wt</i>							
APX	8.67 a	8.62 a	6.17 b	3.20 c	8.91 a	7.49 b	6.59 b	4.49 c
DR 0.67 a	0.77 b	0.70 ab	0.55 c	0.17 a	0.13 ab	0.09 b	0.03 c	
GR 0.41 a	0.58 b	0.48 a	0.33 c	0.62 a	0.60 a	0.42 b	0.36 b	
MR	3.20 a	3.30 a	2.62 ab	1.83 b	1.65 a	1.52 a	1.42 a	1.00 b
Catalase	1340 a	1390 a	1340 a	640 b	160 a	160 a	210 b	230 c

^aStatistical analysis of means (*n*= 6-8) was performed as for Table I.

Table IV. *Antioxidant metabolites and nucleotides in senescing legume nodules*

Metabolite ^a	Bean nodules (days of nitrate)				Pea nodules (days of darkness)			
	0	1	2	4	0	1	2	4
	<i>nmol g⁻¹ fresh wt</i>							
Ascorbate	840 a	1010 b	950 b	840 a	460 a	340 b	300 b	210 c
GSH ^b	103 a	65 b	75 b	37 c	698 a	570 a	378 b	58 c
hGSH ^b	262 a	224 ab	187 b	40 c	90 a	106 a	110 a	36 b
Oxidized thiols ^c	2 a	1 a	2 a	5 b	16 a	14 a	20 b	19 b
NAD ⁺	32.3 a	29.4 ab	26.1 b	17.8 c	29.2 a	18.8 b	13.6 bc	10.2 c
NADH	3.6 a	2.9 b	2.5 bc	2.2 c	11.3 a	7.1 b	4.3 bc	3.3 c
NADP ⁺	9.3 a	7.6 b	6.7 b	4.1 c	4.7 a	4.6 a	2.9 b	2.5 b
NADPH	5.0 a	4.6 a	4.6 a	4.9 a	4.9 a	3.9 ab	3.7 b	3.3 b

^aStatistical analysis of means ($n= 4-12$) was performed as for Table I. ^bDetermined by the HPLC method. ^cThe percentage of thiol tripeptides in the disulfide form was determined by the enzymatic method.

Table V. *Activities of enzymes involved in GSH and hGSH synthesis in senescing legume nodules*

Enzyme ^a	Bean nodules (days of nitrate)			Pea nodules (days of darkness)		
	0	2	4	0	2	4
	<i>nmol min⁻¹ g⁻¹ fresh wt</i>					
γ ECS	5.5 a	3.3 b	2.3 c	4.9 a	5.5 a	7.6 b
GSHS	1.0 a	0.4 b	0.4 b	18.0 a	11.1 b	9.3 c
hGSHS	9.8 a	9.6 a	5.0 b	12.0 a	7.1 b	4.5 c

^aStatistical analysis of means ($n=3-4$) was performed as for Table I.

Table VI. *Oxidant damage of lipids and proteins in senescing legume nodules*

Parameter ^a	Bean nodules (days of nitrate)				Pea nodules (days of darkness)			
	0	1	2	4	0	1	2	4
Lipid peroxides	0.27 a	0.27 a	0.16 b	0.14 b	1.66 a	1.16 b	0.88 b	1.02 b
Oxidized proteins	16.6 a	16.4 a	18.6 a	22.3 b	12.8 a	15.3 ab	17.3 b	16.7 b

^aStatistical analysis of means ($n= 4-8$) was performed as for Table I. Lipid peroxides are expressed in nmol malondialdehyde mg^{-1} lipid and oxidized proteins in nmol carbonyl groups mg^{-1} protein.