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Reduced carbon availability to bacteroids and elevated ureides in nodules, but not in shoots, are involved in the nitrogen fixation response to early drought in soybean

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Nitrogen fixation (NF) in soybean (Glycine max (L.) Merr.) is highly sensitive to soil drying. This sensitivity has been related to an accumulation of nitrogen compounds, either in shoots or nodules, and a nodular carbon flux shortage under drought. In order to assess the relative importance of carbon and nitrogen status on NF regulation, the responses to the early stages of drought were monitored with two soybean cultivars with known contrasting tolerance to drought. In the sensitive cultivar (Biloxi) NF inhibition occurred earlier and was more dramatic than in the tolerant cultivar (Jackson). The carbon flux to bacteroids was also more affected in Biloxi than in Jackson, due to an earlier inhibition of sucrose synthase activity and a larger decrease of malate concentration in the former. Drought provoked ureide accumulation in nodules of both cultivars, but this accumulation was higher and occurred earlier in Biloxi. However, at this early stage of drought, there was no accumulation of ureides in the leaves of either cultivar. These results indicate that a combination of both reduced carbon flux and nitrogen accumulation in nodules, but not in shoots, are involved in the inhibition of NF in soybean under early drought.

Drought-related inhibition of nitrogen fixation (NF) seriously limits legume yield in many arid and semiarid regions of the world. Three major factors have been proposed to be involved in drought effects on NF: oxygen limitation, carbon shortage, and regulation by nitrogen metabolism. The role of oxygen limitation in the response of nitrogenase activity to drought has been discussed extensively (Díaz del Castillo and Layzell, 1995; Serraj and Sinclair, 1996a; Minchin, 1997). However, while it is clear that drought causes an increase in nodular oxygen diffusion resistance (Durand et al., 1987), it is unlikely that such changes are the only cause of the decline in NF, since the inhibition of NF by drought cannot be reverted by simply increasing the O₂ concentration around the rhizosphere (Díaz del Castillo et al., 1994).

An alternative explanation for the decrease in NF under drought is a reduced carbon supply to bacteroids (Arrese-Igor et al., 1999). The main carbon source transported from shoots into nodules is sucrose, which may be hydrolyzed by either sucrose synthase (SS) or alkaline invertase (AI) and studies with rug4 mutants of pea (Pisum sativum L.) have shown that SS activity is essential for nodule functioning (Gordon et al., 1999). SS has been shown to be the first nodule enzyme activity that declines under drought in both tropical legumes, such as soybean (González et al.,
1995) and in temperate legumes such as pea (González et al., 1998; Gálvez et al., 2005). The decrease in SS activity results in an accumulation of sucrose and a reduced concentration of organic acids, mainly malate, in pea nodules (Gálvez et al., 2005), causing a shortage of substrates for bacteroid respiration. As a consequence, a transient accumulation of oxygen in the infected region would take place leading to an increase in the resistance of the oxygen diffusion barrier in order to avoid nitrogenase damage. Both the depletion of respiratory substrates and consequent closure of the oxygen diffusion barrier would cause the observed decline in NF (González et al., 2001). However, this major role of SS for nodule functioning and the response to drought has been recently challenged for the pasture legumes Lotus japonicus L. (Horst et al., 2007) and Medicago sativa L. (Naya et al., 2007).

Nitrogen metabolism has also been proposed to be involved in the regulation of NF under drought conditions by a N feedback mechanism involving shoot N status, with several molecules suggested to be involved in such a mechanism (see King and Purcell, 2005 and references therein). Moreover, it has been reported that NF in ureide-exporter legumes is a more drought-sensitive process than in amide-exporter ones (Sinclair and Serraj, 1995). The ureides, allantoin and allantoate, are the main nitrogenous compounds exported from soybean nodules to shoots (Herridge, 1982) where they are metabolized. An accumulation of ureides under drought in both shoots (Serraj and Sinclair, 1996b; Serraj et al., 1999b) and nodules (Serraj et al., 1999b; Vadez et al., 2000) has been reported. Serraj et al. (2001) suggested that ureide inhibition of nodular activity could occur as either a direct feedback within the nodule or an indirect feedback originated from shoots. The latter has been suggested to depend on ureide catabolism pathways (Sinclair et al., 2003). Allantoin is first metabolised into allantoate by allantoinase and the subsequent step of allantoate degradation in soybean has been attributed to two enzymatic pathways: allantoate amidinohydrolase (Shelp and Ireland, 1985) and allantoate amidohydrolase (Winkler et al., 1985), which requires manganese as a cofactor (Lukaszewski et al., 1992). Moreover, supply of external ureides increased ureides concentration in leaves and inhibited nitrogenase activity (Serraj et al., 1999b; Vadez et al., 2000). Also, Vadez and Sinclair (2001) reported an inverse relationship between shoot ureides concentrations in well-watered plants and the drought sensitivity of NF among nine soybean cultivars, supporting the role of elevated shoot ureides in NF inhibition, although there was no strict correlation between drought sensitivity itself and shoot ureide concentration under drought conditions.

In this study two soybean genotypes, which show different drought sensitivity, have been analyzed. Jackson was identified as having substantial drought tolerance
(Sall and Sinclair, 1991), and subsequently, its NF performance under drought was confirmed under both controlled (Serraj and Sinclair, 1996b; Purcell et al., 1997) and field conditions (Serraj et al., 1997). This drought tolerance was associated with low concentrations of ureides in shoots under drought (Serraj et al., 1997; Serraj and Sinclair, 1997). Biloxi is sensitive to soil drying (Serraj and Sinclair, 1996b) and under drought there is a higher accumulation of ureides in shoots compared to Jackson (Serraj and Sinclair, 1996b; Purcell et al., 1998). This differential accumulation of ureides has been associated with distinct ureide catabolism pathways in leaves (Vadez and Sinclair, 2002), supporting the idea of a systemic regulation of NF. However, Todd and Polacco (2004) have shown that soybean cultivars with contrasting tolerance to drought, such as Mapple Arrow and Williams, use the same enzymatic pathway for allantoate degradation in shoots. Moreover, King and Purcell (2005) recently showed that ureide catabolism in Jackson is not strictly Mn-independent, as earlier proposed. Furthermore, so far there is no available information about nodule carbon fluxes in soybean cultivars showing contrasting tolerance to drought and how these carbon fluxes are correlated with the nitrogen metabolism response at the nodule level. The aim of this study is to characterize the carbon and nitrogen nodule metabolism of these two soybean cultivars at early drought stages to ascertain the relevance of nodule carbon metabolism in NF tolerance to drought.

RESULTS

Drought effects on nodule water status, plant biomass and nitrogen fixation

Plants of both cultivars irrigated at field capacity maintained a similar nodule water potential of -0.50 to -0.60 MPa throughout the study period. However, drought provoked a gradual and progressive decline in the nodule water potential of both cultivars (Fig. 1), with stressed plants showing significant differences from their respective controls 2 days after starting the stress treatment. At the end of the study period, stressed plants of both cultivars reached nodule water potential values of ca. -1.5 MPa (Fig. 1). Plant biomass was similar for both cultivars, with values around 8 g DW plant⁻¹ at the beginning of the experimental period. The mild drought conditions reached in this study did not provoke significant biomass differences between control and stressed plants in either of the cultivars (data not shown).

Specific NF rates, measured as the apparent release of H₂ (ANA) on a nodule biomass basis, were slightly lower in Biloxi than in Jackson under unrestricted water availability. In Jackson, drought provoked a 30% decrease of ANA 3 days after the onset of drought, maintaining this level of activity throughout the study period (Fig. 2A). In contrast, Biloxi showed a 75% ANA decrease within 2 days of the onset of drought.
The protein content of the nodule plant fraction showed similar values for both cultivars, with no differences between well-watered and stressed plants in either of them (data not shown). This suggests that drought level was not sufficient to trigger major, irreversible changes in nodule status.

Drought effects on ureide content of leaves and nodules

In Jackson the ureide content in leaves was around 9 µmol g⁻¹ DW in control plants (Fig. 3A) which was lower than the level in Biloxi (12 µmol g⁻¹ DW, Fig. 3B). Drought did not significantly modify shoot ureide levels in either cultivar throughout the study period (Fig. 3A, B). The nodular ureide content in well-watered Jackson plants was around 35 µmol g⁻¹ NDW and was higher than that in Biloxi nodules (24 µmol g⁻¹ NDW). Drought caused a 2-fold accumulation of ureides in Jackson nodules after 3 days of treatment which reached a 2.5-fold accumulation at the end of the study (Fig. 3C). In contrast, Biloxi nodules showed a 2-fold ureide accumulation 2 days after the onset of drought, which increased to a 5-fold accumulation at the end of the drought period (Fig. 3D).

Drought effects on nodular enzyme activities and organic acids content

Regarding the enzyme activities monitored, the most immediate and marked response to drought was that observed for SS (Fig. 4A, B). Under unlimited water supply, nodule SS activity was slightly, although significantly, higher in Jackson than in Biloxi (0.174 ± 0.009 vs 0.151 ± 0.005 µmol NADH mg⁻¹ protein min⁻¹). In the sensitive cultivar Biloxi, SS activity showed a 25% decrease within 1 day of water deprivation, declining to 55% at the end of the study (Fig. 4B). However, in Jackson, a significant decrease of this activity was not observed until 3 days after the onset of drought and was only reduced by 40% at the end of the study period (Fig. 4A). To test whether this decline of activity was due to a reduction of protein content, an immunodetection assay was carried out (Fig. 4C). SS immunodetection showed a reduction in the protein amount similar for both cultivars.

Both cultivars showed similar values for alkaline invertase (AI), malate dehydrogenase (MDH), NADP⁺-dependent isocitrate dehydrogenase (ICDH), aspartate aminotransferase (AAT) and glutamate synthase (GOGAT) activities (average value for control nodules of both cultivars were 0.18, 8.5, 0.08, 0.5, and 0.026 µmol mg⁻¹ protein min⁻¹, respectively). Only the activity of PEPC was markedly higher in Jackson (0.14 ± 0.004 µmol mg⁻¹ protein min⁻¹) than in Biloxi (0.06 ± 0.004 µmol mg⁻¹ protein min⁻¹). In Jackson, none of these activities was significantly affected by drought, whereas in Biloxi AI, AAT and GOGAT activities decreased by 25%, at the end of the treatment period (data not shown).
Nodular malate content in Jackson control plants was 30.2 ± 1.2 µmol g⁻¹ NDW, and was 2.5-fold higher than in control plants nodules of Biloxi (Fig. 5). Succinate concentrations in Jackson nodules were also significantly higher than those of Biloxi, although succinate concentrations were much lower than those of malate (1.25 ± 0.08 and 0.31 ± 0.03 µmol g⁻¹ NDW in Jackson and Biloxi, respectively). Citrate and 2-ketoglutarate contents did not show significant differences between cultivars (data not shown). Malate decreased significantly in both cultivars 2 days after the onset of drought. However, the decline of malate content was greater in Biloxi as nodule water potential became more negative (Fig. 5). At the end of the study period Jackson nodules exhibited malate levels ca. 70% of those of control plants, whereas in Biloxi nodules the levels were reduced to 50% of control values (Fig. 5).

Table 1 shows the linear regressions between ANA and ureide concentration, malate concentration and the ratio of malate/ureide concentrations, respectively, in nodules of stressed plants. Regressions were obtained using values previously normalized against their respective controls for each cultivar. For ureides, the correlation coefficients (r² values) for both cultivars are very similar, at around 0.5. However, the correlation of ANA with malate increased to a range of 0.7-0. and was even higher with the malate/ureides ratio, with values above 0.9 for both cultivars. Indeed, when regression analysis was performed including both cultivars, the correlation between normalized ANA and normalized malate/ureides ratio remained 0.76, despite the relative different response of each cultivar to drought.

DISCUSSION

Nodule water status and nitrogen fixation

There is a high diversity among the response of different legume species to water stress (Sinclair and Serraj, 1995). Indeed, such diversity can be found within different lines of the same species, i.e. different soybean cultivars showing distinct tolerances or sensitivities to drought (Sall and Sinclair, 1991; Serraj and Sinclair, 1997). In this study drought provoked a similar nodule water potential decline in both Jackson and Biloxi soybeans (Fig. 1). Most published papers linking drought and ureide effects on NF express water stress as either fractions of transpirable soil water, a parameter of relevant agronomical significance, or leaf water potential, which is the usual way to express plant water status. Less attention has been given to nodule water potential which, although closely related to leaf water potential (González et al., 1995), has been recently shown to be more relevant to NF than plant (leaf) water status (Marino et al., 2007a).
Drought inhibited NF in both Jackson and Biloxi, but this inhibition occurred earlier and more severely in Biloxi (Fig. 2). These results, measured by a flow-through gas system detecting H₂ evolution, confirm that NF in Jackson is more tolerant to drought, as already reported in field and greenhouse experiments, using acetylene reduction techniques (Serraj and Sinclair, 1996b; Serraj et al., 1997).

Ureides accumulate in nodules, but not in leaves, in early drought

Studies of Sinclair’s group established a relationship between the greater tolerance to drought of Jackson and a smaller ureide accumulation in leaves (Serraj and Sinclair, 1997; Serraj et al., 1997; Vadez and Sinclair, 2002). Indeed, Sinclair et al. (2003) suggested that the differences in leaf ureide metabolism was the main cause of the distinct sensitivity of the cultivars to drought. In general, these studies were carried out over medium-term periods of drought or following the supply of external ureides. It can be argued that experiments using an external supply of ureides, in order to mimic the physiological response to drought, do not allow these ureides to reach the nodules (the site of their physiological synthesis) via the xylem. Instead, they are directly targeted to leaves. This would then preclude the possibility of a build-up of these compounds in nodules, and therefore would not reflect the natural physiological consequences of a drought-induced impairment of long-distance transport. Furthermore, King and Purcell (2005) suggested that shoot ureide accumulation in soybean is not responsible for NF inhibition. In the present study, no accumulation of ureides in leaves of Jackson or Biloxi has been detected (Fig. 3A, B) despite the fact that NF was already inhibited (Fig. 2A, B). Taken together, these results indicate that leaf ureides are not involved at the early stages of NF inhibition under drought, although a role in later stages of a more severe drought (Serraj et al., 1999b) can not be discarded.

A possible role of nodule ureide content in soybean NF has received much less attention, despite the fact that more than 30 years ago Minchin and Pate (1974) recorded an increase in the soluble amino acid content of pea nodules under low transpiration conditions. There is only one very recent comparative study on nodule ureide content profiles between drought-tolerant and -sensitive cultivars (King and Purcell, 2005), which showed that nodular ureide accumulation mirrored the decline in NF. Therefore, these authors concluded that ureides, along with other N-compounds, are potential candidates for inducing a feedback inhibition of NF. In the present work, drought provoked ureide accumulation in nodules of both cultivars. As this accumulation was greater and occurred earlier in Biloxi (Fig. 3C, D), it appears to be correlated with the NF sensitivity to drought of this cultivar. It is noteworthy that ureide accumulation occurs in nodules despite a decreased NF. Although some other
causative factors may also lead to this effect, such as an increase in the activity of ureide biosynthetic enzymes, this seems to unlikely since, based on our previous experience, nodular metabolic activities of primary metabolism tend to be stable or to decline under mild or moderate drought (Larrainzar et al., 2007). The only exception to this trend is ICDH, whose activity has been shown to slightly increase in pea nodules both under drought (Gálvez et al., 2005) and oxidative stress (Marino et al., 2007b). Thus, it can be concluded that export is impaired, resulting in the accumulation of certain metabolites. The circumstances leading to this impaired export have been discussed elsewhere (Streeter, 1993; Walsh, 1995; Serraj et al., 1999a). Our results do not provide evidence for nodule ureides being the actual compounds which induce the decrease in NF, but it is likely that an accumulation of fixation products is involved. Indeed, Lodwig et al. (2003) have shown that a complex amino-acid cycling occurs in nodules, with the plant providing amino acids to the bacteroids, enabling them to shut down their ammonium assimilation, and bacteroids acting like plant organelles to cycle amino acids back to the plant for asparagine synthesis. However, it is not yet known if this exchange could be directly disrupted by a drought-induced accumulation of N-compounds.

Furthermore, the present results are also in agreement with the hypothesis that the cause of NF inhibition under drought is of a local origin, rather than relying on a systemic signal (Marino et al., 2007a).

Carbon flux in nodules is more affected in the sensitive cultivar

SS has already been described as a key enzyme which regulates NF under drought conditions in soybean (González et al., 1995) and pea (González et al., 1998; Gálvez et al., 2005). The activity of this enzyme has also been observed to decline under other abiotic stresses such as salinity, defoliation, nitrate and oxidative stress (Fernández-Pascual et al., 1996; Gordon et al., 1997; Marino et al., 2006). Moreover, there is a high correlation between SS inhibition and the NF decline under abiotic stresses (Arrese-Igor et al., 1999). However, it has yet to be proven that diversity in SS activity can be associated with a differential tolerance to drought. In the present work, SS was the first nodule enzyme to show reduced activity under drought in both cultivars (Fig. 4A, B), with the inhibition occurring at the first day of reduced watering in Biloxi (Fig. 4B), and before any observed effect on NF (Fig. 2B). In contrast, Jackson maintained SS activity rates at control values until the third day of drought (Fig. 4A), and the decline was concomitant with that of NF (Fig. 2A). The SS activity decline was also related to a reduction in SS content (Fig. 4C). However, in Biloxi the activity decline was observed prior to any effect on SS protein content, suggesting that SS
activity could be modulated through post-transcriptional modifications (Komina et al., 2002; Geigenberger, 2003).

In pea nodules, a decreased SS activity leads to a decline in malate content (Gálvez et al., 2005; Marino et al., 2007a). It had been earlier suggested that a likely decrease in nodule malate content under certain environmental constraints may lead to the inhibition of NF (Arrese-Igor et al., 1999). In the present work, malate content in Jackson nodules ranged between 2- and 3-fold the measured values in Biloxi nodules (Fig. 5), a situation that is consistent with the higher NF rates of the former and seems to be related to the higher SS and, particularly, PEPC activity rates. A decline in nodular malate content was observed after 2 days of drought treatment in both cultivars, but it was more pronounced in Biloxi (Fig. 5), a fact that is not related to PEPC, whose activity was maintained throughout the study period in both cultivars at control levels, but instead to the decline of SS activity. The higher constitutive content of malate may also be an important component of the drought tolerance of Jackson nodules, since a greater malate availability would reduce the effect of a carbon-skeleton shortage due to SS down-regulation.

**Carbon/nitrogen interactions in the inhibition of nitrogen fixation under drought in soybean**

In order to determine the relative importance of C or N in the inhibition of NF under drought, correlation analyses were performed between normalized ANA and normalized ureide content, malate content and the malate/ureide ratio (Table 1) in nodules of stressed plants. The correlation between malate and ANA was higher than that of ureides and ANA reflecting the involvement of carbon shortage in the inhibition of NF during early drought conditions. However, when both factors were combined, NF dependence on the malate/ureide ratio showed a higher correlation than that of ureides or malate alone. A strong relation between C and N metabolism is widely accepted as a crucial interplay in the regulation of plant performance and it has also been shown to occur in water-stressed pea nodules (Gálvez et al., 2005). Taken together, these results indicate that there are at least two major mechanisms involved in the drought-induced inhibition of NF in soybean nodules: 1) an impairment of long distance transport, leading to an accumulation of ureides in nodules, which is likely to provoke a N feedback regulation of NF; 2) an impairment of metabolic C flux in nodules, resulting in a shortage of C substrate for bacteroid NF. Both factors seem to be crucial for the regulation of NF under drought, although the results obtained so far do not allow for a definitive statement such as whether the accumulation of N-compounds has a direct effect on NF or an indirect one through an induced decline in SS activity. Furthermore,
both N accumulation and SS decline could be independently induced by oxidative signaling. This has been recently suggested for pea (Marino et al., 2006) and alfalfa (Naya et al., 2007) and is consistent with the regulation of NF under drought occurring at the local/nodule level (Marino et al., 2007a), rather than systemically.

MATERIALS AND METHODS

Growth conditions

Soybean plants (Glycine max (L.) Merr.) cvs Jackson and Biloxi were inoculated with the hup’ Bradyrhizobium japonicum strain UPM792, to allow for the detection of H₂ evolution. Plants were grown in 1 L pots with a 2/1 (v/v) mixture of vermiculite/perlite as rooting substrate in a controlled environmental chamber (24/18 °C day/night temperature, 60/70% day/night relative humidity and 16 h photoperiod). They were watered three times a week with nutrient solution lacking N (Rigaud and Puppo, 1975).

Experimental procedures, nitrogen fixation and water potential

In order to obtain plants with similar biomass and developmental stage, experiments were carried out when plants were 5 and 6 weeks old for Biloxi and Jackson cultivars, respectively. Previous experiments using 6 weeks old Biloxi plants gave an identical similar metabolic profiling, but, due to their bigger size, transpiration rates were higher and the water content of pots was more rapidly depleted. Therefore, when analyzed under these conditions, drought effects were more dramatic and less comparable between cultivars. Plants were separated randomly into two sets: control and drought. During the study period, control plants were supplied daily with nutrient solution to field capacity, whereas stressed plants were supplied daily with 1/4 of the measured evapotranspirational water loss volume. Four plants per treatment were harvested at days 1, 2, 3 and 6 after the onset of drought in order to obtain data at different levels of stress.

For apparent nitrogenase activity (ANA) determinations, H₂ evolution of intact plants, whose root systems were sealed into the growth pots, was measured in an open flow-through system under N₂/O₂ (79%/21%) according to Witty and Minchin (1998) using an electrochemical H₂-sensor (Qubit System Inc., Canada). The H₂ sensor was calibrated using high purity gases (Praxair, Madrid, Spain) employing a gas mixer (Air Liquide, Madrid, Spain) flowing at the same rate as the sampling system (500 mL min⁻¹).

Nodule water potential was determined by a Wescor HR-33T psychrometer (Wescor Inc. 5500, Logan, UT, USA). Nodules were harvested, frozen in liquid N₂ and
stored at –80°C for further analysis. Roots and shoots were separated and dried for 48 h at 70°C for dry weight determinations.

**Extraction and assay of enzymes**

All enzymes were extracted from nodules at 4°C with mortar and pestle in an optimized medium consisting of 50 mM MOPS, pH 7, 20% PVPP, 10 mM DTT, 10 mM 2-mercaptoethanol, 1 mM EDTA, 20 mM KCl and 5 mM MgCl₂ (Marino et al., 2006). The homogenates were centrifuged for 30 min at 20 000 g, 4 ºC. Aliquots of the supernatants were retained for determination of plant fraction protein (Bradford, 1976) and for PEPC (EC 4.1.1.31) activity. The rest of the supernatant was desalted by low-speed centrifugation (180 g, 2 min) through 5-mL columns of Bio Gel P6DG (BioRad) previously equilibrated with 50 mM MOPS pH 7, 20 mM KCl, 5 mM MgCl₂. Desalted extracts were used to assay SS (EC 2.4.1.13), Al (EC 3.2.1.26), MDH (EC 1.1.1.37), GOGAT (EC 1.4.1.14), AAT (EC 2.6.1.1) and ICDH (EC 1.1.1.42). All activities were measured within the linear range at 30ºC, as described elsewhere (Marino et al., 2006).

**Malate and ureides determination**

To analyze malate and ureide content of nodules, frozen nodules were homogenized to a fine powder in liquid N₂ with mortar and pestle. 1.5 mL of 10 % (w/v) TCA in water was then added and the homogenate was centrifuged for 10 min at 1 750 g, 4°C. The aqueous phase was washed six times with diethyl ether saturated with water. The diethyl ether was discarded and the aqueous phase was purged with He for two min and then filtered through a 0.45 µm syringe filter (Wilson and Harris, 1966).

Malate levels were determined by ion chromatography in a DX-500 system (Dionex) by gradient separation with a Dionex IonPac AS11 column according to the manufacturer's instructions (2.5 mM NaOH/18 % methanol to 45 mM NaOH/18 % methanol in 13 min (Gálvez et al., 2005).

Nodule ureides (allantoin and allantoate) were determined by capillary electrophoresis. The length of the capillary tube was 60 cm and 0.1 M Na₂B₄O₇•10H₂O (pH 9.2), 25 mL L⁻¹ OFM-Anion BT (Waters) solution was used as electrolyte. Samples were injected for 5 s by the hydrostatic method, and electrophoresed under 10 kV for 30 min. Allantoin and allantoate were detected by optical density at 190 nm (Sato et al., 1998).

Ureide content of leaves was analyzed following extraction in 1 mL of 0.2 N NaOH to 10 mg of dry tissue, boiling of extracts for 30 min and centrifugation at 12 000 g for
10 min. Ureides were quantified using a colorimetric detection method (Trijbels and Vogel, 1966).

**Immunoblotting**

SDS-PAGE was performed according to Laemmli (1970) with a 1-mm thick 10% polyacrylamide (w/v) resolving gel and a 4.6% (w/v) stacking gel in a vertical electrophoresis cell (MiniProtean III, Bio-Rad, Hercules, CA, USA) at 150 V for 60 min. Gels were electroblotted onto PVDF membranes for 75 min at 100 V in a Mini Trans-Blot Electrophoretic Transfer Cell (Bio-Rad, Hercules, CA, USA). Blots were blocked in 5% (w/v) skim milk in 20 mM Tris-buffered saline at 4°C overnight. As primary antibody anti-SS (1:5000, v/v) was used. As the secondary antibody goat anti-rabbit IgG alkaline phosphatase (1:10.000, v/v Sigma-Aldrich, St. Louis, USA) was employed. Immunoreactive bands were visualized with a bCIP/NBT liquid substrate system (Sigma-Aldrich, St. Louis, USA).

**Statistical analysis**

Results were examined by two-ways analysis of variance. All effects discussed in this study were significant at P ≤ 0.05 in Fisher’s least significant difference (LSD) among means.

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Table 1. Regression line equations between normalized ANA (y) and normalized ureide content, malate content and the ratio between malate and ureide content of Jackson and Biloxi nodules (x in each column, respectively). Each equation is followed by its correlation coefficient (r²). Values were obtained from those given in figures 2-5, normalized against their respective daily control values.

| Cultivar | Ureides  | Malate    | Malate / Ureides |
|----------|----------|-----------|------------------|
| Jackson  | y = -0.132 x + 0.895 | y = 0.591 x + 0.258 | y = 0.261 x + 0.570 |
|          | r² = 0.50 | r² = 0.77 | r² = 0.94 |
| Biloxi   | y = -0.129 x + 0.781 | y = 1.179 x - 0.384 | y = 0.639 x + 0.155 |
|          | r² = 0.52 | r² = 0.70 | r² = 0.90 |
Figure legends

**Figure 1.** Effects of drought on nodule water potential of Jackson and Biloxi soybean plants. For Jackson an asterisk (*), and for Biloxi a hash (#) represent significant differences with the corresponding control values at $P \leq 0.05$. Values represent mean ± standard error ($n = 4$).

**Figure 2.** Effects of drought on apparent nitrogenase activity (ANA) of Jackson (A) and Biloxi (B). Legends and statistical analysis are as described in Figure 1. NDW denotes nodule dry weight. Values represent mean ± standard error ($n = 4$).

**Figure 3.** Effects of drought on ureides levels (allantoin and allantoic acid) of Jackson (A) and Biloxi (B) leaves and Jackson (C) and Biloxi (D) nodules. Legends and statistical analysis are as described in Figure 1. LDW and NDW denote leaf and nodule dry weight, respectively. Values represent mean ± standard error ($n = 4$).

**Figure 4.** Effects of drought on sucrose synthase activity (SS) of Jackson (A) and Biloxi (B) nodules. Legends and statistical analysis are as described in Figure 1. Values represent mean ± standard error ($n = 4$). Panel C shows western immunoblots of host plant SS levels from control (C) and stressed (D) plants, at days 0, 1, 2, 3 and 6 after the onset of drought. Equal amounts of protein were loaded on each lane. Densitometry analysis displayed, expressed as a percentage of the control values, corresponds to four independent biological experiments.

**Figure 5.** Effects of drought on malate levels of Jackson and Biloxi nodules. Legends and statistical analysis are as described in Figure 1. NDW denotes nodule dry weight. Values represent mean ± standard error ($n = 4$).
Figure 1. Ladrera et al. 2007
Figure 2. Ladrera et al. 2007
Figure 3. Ladrera et al. 2007
Figure 4. Ladrera et al. 2007
Figure 5. Ladrera et al. 2007

Malate (µmol g\(^{-1}\) NDW) vs Time (days)