Effect of Changes in Shoot Carbon-Exchange Rate on Soybean Root Nodule Activity¹

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ABSTRACT

The effect of short- and long-term changes in shoot carbon-exchange rate (CER) on soybean (Glycine max [L.] Merr.) root nodule activity was assessed to determine whether increases in photosynthate production produce a direct enhancement of symbiotic N₂ fixation. Shoot CER, root + nodule respiration, and apparent N₂ fixation (acetylene reduction) were measured on intact soybean plants grown at 700 microeinsteins per meter per second, with constant root temperature and a 14/10-hour light/dark cycle. There was no diurnal variation of root + nodule respiration or apparent N₂ fixation in plants assayed weekly from 14 to 43 days after planting. However, if plants remained in darkness following their normal dark period, a significant decline in apparent N2 fixation was measured within 4 hours, and decreasing CO₂ concentration from 320 to 90 microliters CO₂ per liter produced diurnal changes in root nodule activity. Increasing shoot CER by 87, 84, and 76% in 2-, 3-, and 4-week-old plants, respectively, by raising the CO₂ concentration around the shoot from 320 to 1,000 microliters CO₂ per liter, had no effect on root + nodule respiration or acetylene-reduction rates during the first 10 hours of the increased CER treatment. When the CO2-enrichment treatment was extended in 3-weekold plants, the only measured parameter that differed significantly after 3 days was shoot CER. After 5 days of continuous CO₂ enrichment, root + nodule respiration and acetylene reduction increased, but such changes reflected an increase in root nodule mass rather than greater specific root nodule activity. The results show that on a 24-hour basis the process of symbiotic N₂ fixation in soybean plants grown under controlled environmental conditions functioned at maximum capacity and was not limited by shoot CER. Whether N₂-fixation capacity was limited by photosynthate movement to root nodules or by saturation of metabolic processes in root nodules is not known.

Symbiotic N₂ fixation in legumes requires carbon substrates derived from photosynthate or from stored reserves (9). The substrates provide a source of energy for N₂ fixation and acceptor molecules for transporting reduced N. The original source of such carbon compounds is primarily photosynthetic CO₂ reduction. Claims that photosynthate limits N₂ fixation in soybean (3-5) were supported by long-term CO₂-enrichment studies in fieldgrown plants. Those experiments showed that CO₂ enrichment increased total N content and promoted both the rate of acetylene reduced/unit nodule mass and the total nodule mass after 7 d (3-5). Such evidence indicates the importance of photosynthate for N₂ fixation, but, without shoot CER³ measurements and information on short-term responses, one cannot determine whether photosynthate production or transport limits N_2 fixation in soybean.

Two types of experiments suggest that photosynthate availability in nodules rather than shoot CER limits N_2 fixation in soybean. Widely different values of soybean shoot CER produced by varying CO₂ concentrations had little effect on the maximum acetylene-reduction rate during a 24-h photoperiod (13). However, because a 40-h dark pretreatment was used to achieve other experimental objectives, it is difficult to extrapolate from those results to normal plants. Grafting techniques that produced soybean plants with 2:1 or 1:1 shoot:root ratios also suggest that photosynthate availability rather than CER limits N_2 fixation, but no direct measurements of shoot CER or photosynthate movement were made (16). Thus, short-term increases in acetylene reduced/ g nodule tissue could have been associated with increased photosynthate transport or with a trauma-induced utilization of root carbohydrates.

The purpose of the present study was to test whether increases in soybean shoot CER of normal plants produce short-term changes in root nodule activity. When it was observed that no significant change in root nodule activity could be detected within 10 h after increasing CER, it became important to determine how long-term changes in shoot CER produce the previously reported (3-5) increase in acetylene reduction after 7 d of CO₂ enrichment. The experiments, therefore, also address the question of how an important agronomic plant responds to an increase in CER and achieves a new equilibrium between CO₂ and N₂ reduction.

MATERIALS AND METHODS

Growth Conditions. 'Clark' soybeans (*Glycine max* [L.] Merr.) were germinated and planted as previously described (18). Plants were grown under controlled conditions at a photon flux density of 700 μ E m⁻² s⁻¹ (400–700 nm), a 14-/10-h light/dark photoperiod at 28/25 °C, and 70% RH. Root temperatures were maintained at 25 ± 1 °C. Carbon dioxide was controlled at either 320 ± 20 or 1,000 ± 75 μ l CO₂/l of air by continuous monitoring with an IR gas analyzer. Plants were inoculated with *Rhizobium japonicum* strain USDA 3IIb110 and watered with a nutrient solution containing 2 mm NO₃⁻ (18). Under these conditions, soybeans flowered approximately 31 days after planting.

Gas-Exchange Measurements. Rates of apparent whole-plant photosynthesis were measured by differential IR gas analysis in Plexiglas chambers (18) with environmental conditions identical to those under which the plants had grown. Root + nodule respiration was measured with the same system by passing air through the rooting medium which was separated from the shoot system at all times by a Plexiglas lid sealed to the pot. Measurements of apparent N₂ fixation were made on intact plants by determining acetylene-dependent ethylene production, less rigorously termed acetylene reduction (18).

Compositional Analyses. Plants were harvested and dried at

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³ Abbreviation: CER, carbon-exchange rate.

70°C for 48 h. Sucrose, fructose, glucose, pinitol, and starch were measured by gas chromatography (13). Values recorded for sucrose, fructose, glucose, and pinitol were summed to estimate total soluble sugar.

RESULTS

Root + nodule respiration and apparent N₂ fixation showed no significant diurnal variation under normal environmental conditions 2 to 6 weeks after planting (Figs. 1 and 2). When the dark period was extended beyond the customary night period, both root + nodule respiration and apparent N₂ fixation declined dramatically, compared with plants exposed to their normal photoperiod (Fig. 3). When plants were grown under normal CO₂ conditions (320 μ l CO₂/l) and then transferred to a low CO₂ concentration (90 μ l CO₂/l), a marked diurnal variation in both root + nodule respiration and apparent N₂ fixation was measured 5 d later (Table I).

Increasing shoot CER by raising the CO₂ concentration from 320 to 1,000 μ l CO₂/l for 10 h produced no significant change in root + nodule respiration or apparent N₂ fixation in plants of five ages (Table II). When plants were grown at the higher CO₂ concentration for longer periods of time, significant differences in respiration and total acetylene reduction between the two treatments were measured after 5 d (Table III), but there was no significant effect of CO₂ enrichment on specific acetylene-reduction activity (defined as μ mol ethylene g⁻¹ nodule dry weight h⁻¹).



FIG. 1. Diurnal time course of root + nodule respiration at intervals up to 6 weeks after planting. Dark periods are indicated by shading on the upper axis. Each point represents the mean \pm sE of four replicate plants. There was no significant diurnal variation of root + nodule respiration in 14-d-old plants.



FIG. 2. Diurnal time course of apparent N₂ fixation at intervals up to 6 weeks after planting. Dark periods are indicated by shading on the upper axis. Each point represents the mean \pm sE of five replicate plants. There was no detectable acetylene-reduction activity in 14-d-old plants.



FIG. 3. Apparent N₂ fixation and root + nodule respiration of 21-dold plants kept in continuous darkness following the normal dark cycle (O, Δ) . Control plants (Φ , \blacktriangle) grown under the normal light/dark cycle (indicated by shading on the upper axis) are included for comparison. Each point represents the mean \pm sE of four plants.

Table I. Diurnal Changes in Root + Nodule Respiration and Apparent N_2 Fixation of Soybeans in a CO_2 -Depleted Atmosphere

Plants were grown at $320 \pm 20 \ \mu l \ CO_2/l$ for 23 d and then maintained at $90 \pm 10 \ \mu l \ CO_2/l$ for 5 d. A 10-h dark period was imposed between 1800 and 0400 h. The rates of root + nodule respiration and apparent N₂ fixation on day 23 for the plants were $8.6 \pm 0.2 \ \text{mg} \ CO_2 \ \text{plant}^{-1} \ h^{-1}$ and $20.4 \pm 1.3 \ \mu\text{mol}$ ethylene plant⁻¹ h⁻¹. Shoot CER on day 28 for control plants and plants transferred to low CO₂ were 71.2 $\pm 2.5 \ \text{and} \ 18.2 \pm 0.9 \ \text{mg} \ CO_2 \ \text{plant}^{-1} \ h^{-1}$, respectively. Root + nodule respiration and apparent N₂ fixation in control plants on day 28 were 14.7 mg CO₂ plant⁻¹ h⁻¹ and 35.7 \ \mu\text{mol} ethylene plant⁻¹ h⁻¹. All values are the mean $\pm \text{se}$ of three replicate plants.

Time	Root + Nodule Respiration	Apparent N ₂ Fixation		
h	mg CO_2 plant ⁻¹ h ⁻¹	μ mol ethylene plant ⁻¹ h ⁻¹		
1730	8.0 ± 1.5	18.5 ± 2.0		
2230	4.7 ± 0.5	13.7 ± 2.0		
0330	4.4 ± 0.5	10.3 ± 2.1		
0830	7.0 ± 0.8	17.6 ± 2.0		

Table II. Shoot CER, Root + Nodule Respiration, and Apparent N_2 Fixation of Soybean Plants before and after 10-Hour Exposure to 1,000 μ l CO₂/l

Plant Age	Shoot CER		Root + nodule Respiration		Apparent N ₂ Fixation	
	Before	After	Before	After	Before	After
days	mg CO_2 plant ⁻¹ h ⁻¹			μ mol ethylene \cdot plant ⁻¹ h ⁻¹		
14	4.4	8.2ª	0.8	0.9	b	
21	13.7	25.4°	3.6	4.4	10.3	9.6
27	55.8	98.3 ^d	8.6	9.0	26.1	28.0
36	e		25.0	26.2	75.9	76.2
43			48.5	49.4	116	131

^a CO₂ treatment effect significant at $p \le 0.001$ (df = 6).

^b There was no detectable acetylene-reduction activity in 14-d-old plants.

^c CO₂ treatment effect significant at $p \le 0.05$ (df = 6).

^d CO₂ treatment effect significant at $p \le 0.01$ (df = 6).

^e CER was not measured after day 27, because plants were too large for the assay chamber.

Table III. Root + Nodule Respiration and Apparent N_2 Fixation ofSoybeans Exposed to a CO2-Enriched Atmosphere

Plants were grown at 320 μ l CO₂/l continuously or transferred to 1,000 μ l CO₂/l 23 d after planting. Values represent the mean of five replicate plants. Shoot CER of soybeans exposed to 1,000 μ l CO₂/l was 73% greater on day 2 and 58% greater on day 5 than was that of plants grown at 320 μ l CO₂/l. Specific activity of apparent N₂ fixation on day 6 was 186 ± 6 and 198 ± 18 μ mol ethylene g⁻¹ nodule dry weight h⁻¹ at 320 and 1,000 μ l CO₂/l, respectively.

Time after Transfer		· Nodule iration	Apparent N ₂ Fixation		
	320	1,000	320	1,000	
	μl CO ₂ /l				
days	mg CO ₂	$plant^{-1}h^{-1}$	µmol ethyler	$ne \cdot plant^{-1} h^{-1}$	
0	9.7	9.6	21.9	19.4	
2	11.2	11.5	26.6	28.3	
3	11.4	13.4	28.4	33.8	
5	14.7	18.1ª	35.7	49.7 ^ь	
6	18.3	23.1 ^b	51.2	66.9 ^b	

* CO₂ treatment effect significant at $p \le 0.01$.

^b CO₂ treatment effect significant at $p \le 0.05$.

Table IV. Dry Weight, Soluble Sugar, and Starch Content of 29-Day-Old Soybeans

Plants were grown continuously at 320 μ l CO₂/l or transferred on day 23 to 1,000 μ l CO₂/l. Values are the mean of five replicate plants on a dryweight basis.

Plant Organ	Dry Weight		Soluble Sugar		Starch	
	320	1,000	320	1,000	320	1,000
	μl CO ₂ /l					
	g/µ	olant		m	g/g	
Leaves	1.76	2.92ª	43.9	43.6	27.9	77.8 ^b
Stem	0.80	0.95 ^b	33.0	32.0	2.4	5.9
Root	1.01	1.16	12.1	15.4	0.9	1.4
Nodules	0.27	0.34 ^b	13.6	10.7	7.4	7.7

^a CO₂ treatment effect significant at $p \le 0.01$.

^b CO₂ treatment effect significant at $p \le 0.05$.

Significant effects of CO_2 on dry matter accumulation of leaves, stems, and nodules were measured after 6 d (Table IV). However, carbohydrate analyses showed that only the leaves of CO_2 -enriched plants had a significantly greater concentration of nonstructural carbohydrates and that the increase resulted from starch accumulation (Table IV).

Thirty-five-d-old soybean plants, pretreated for 2 d with 1,000 μ l CO₂/l and then kept in darkness following the normal dark cycle, maintained acetylene reduction equal to or greater than initial rates for 19 h after the lights normally would have come on (Fig. 4). Control plants not exposed to CO₂ enrichment responded in a manner similar to that of younger plants, with apparent N₂ fixation declining soon after lights normally would have come on (Fig. 4 *versus* Fig. 3). In contrast, plants that were grown with 1,000 μ l CO₂/l for 28 d maintained normal acetylene-reduction rates for only 5 h during an extension of the normal dark period (Fig. 5).

DISCUSSION

Results from this study clearly demonstrate that soybean root nodules on intact, growing plants can reduce acetylene at a stable rate without diurnal fluctuations. That conclusion is consistent with studies of soybeans examined at one stage of development



FIG. 4. Apparent N₂ fixation of 35-d-old soybean plants during an extended dark treatment following the normal night period. Plants were either grown at 1,000 μ l CO₂/l for 2 d before the initiation of the experiment at time 0 (**•**) or kept at 320 μ l CO₂/l throughout (\bigcirc). The first value represents the mean ± sE of six plants (three from each treatment) taken 30 min before the growth chamber lights were to have come on. Subsequent values represent the mean ± sE of five replicate plants.



FIG. 5. Apparent N₂ fixation of 28-d-old soybean plants during an extended dark treatment following the normal night period. Plants were grown continuously in 1,000 μ l CO₂/1 (\odot) or in 320 μ l CO₂/1 (\bigcirc) from time of planting. Time 0 represents the end of the normal night period. Each point represents the mean ± sE of five replicate plants. Specific activities of acetylene reduction for high and control CO₂ treatments at the start of the experiment were 223 ± 13 and 238 ± 27 μ mol ethylene g⁻¹ nodule dry weight h⁻¹, respectively.

under constant temperatures (2, 12) but contrasts with observations by other workers when diurnal temperatures varied. Neither root + nodule respiration nor acetylene reduction showed any significant decline during the night period in plants assayed weekly between 14 and 43 d after planting (Figs. 1 and 2). Such fluctuations would be expected if carbon substrates available to roots and nodules declined below a critical level at any time during each 24h period. That assumption is supported by the observation that acetylene reduction and root + nodule respiration declined significantly when the normal dark period was extended (Fig. 3) and by the demonstration that those parameters declined significantly during a normal night period when plants were shifted from 320 to 90 μ l CO₂/1 for 5 d (Table I). Such data support the concept that the nodule carbohydrate pool is limited but do not indicate whether an increase in CER could enhance N₂ fixation.

Increasing CER by providing 1,000 rather than 320 μ l CO₂/l had long-term, but not short-term, effects on root nodule activity. Significant amounts of new photosynthate move to soybean roots within 1 h after exposure to ¹⁴CO₂ (6, 11), and, even in carbohy-

drate-depleted soybeans, maximum rates of acetylene reduction were evident after fewer than 4 h in the light (13). Nevertheless, large increases in CER produced by CO₂ enrichment of air surrounding shoots of plants of five ages, including two measurements made after anthesis, had no significant promotive effect on root + nodule respiration or acetylene reduction within 10 h (Table II). When the CO₂-enrichment treatment was extended, however, significant increases in root + nodule respiration and acetylene reduction were measured after 5 d (Table III). Unlike previously reported studies (3-5), the increase in acetylene reduction after 6 d of CO₂ enrichment reflected primarily nodule growth (Tables III and IV). Although Sheehy et al. (14) found a significant positive correlation between whole plant CER and acetylene reduction, those alfalfa plants with greater CER values also had greater plant and root nodule dry weights. In the present experiments, no difference was measured in acetylene reduction values between control and experimental plants until plants with greater CER values had significantly greater nodule dry weight. The slight increase in acetylene-reduction rate/g nodule dry weight associated with CO₂ enrichment (Tables III and IV) might have developed into a significant difference later and, therefore, may be comparable to previous results (3-5). These data emphasize that root nodules of soybean plants do not always have excess acetylene-reduction capacity that can be activated by CER values greater than those typically experienced by the plants. Whether the situation reported after 7 d of CO₂ enrichment for field-grown soybeans (3-5) actually reflected an excess of acetylene-reduction capacity on day 0 can be determined only by the type of data in Table III.

Root nodule activities during an extended dark period also differed greatly between plants given short- or long-term CO₂ enrichment treatments. Plants exposed to 1,000 µl CO₂/l for 2 d showed no decline in acetylene-reduction activity of nodules during the first 19 h of an extended dark treatment following the normal night period (Fig. 4). When plants were grown for 28 d in 320 or 1,000 μ l CO₂/l treatments, acetylene-reduction activity declined significantly during 11 h of a similar dark treatment (Fig. 5). The latter type of results have been used to indicate that plants grown under constant environmental conditions were in a steady state of substrate production and use on a 24-h basis (7). Although the total acetylene-reduction activity of plants exposed to the higher CO₂ level for 28 d was nearly twice that of plants grown with a normal CO₂ concentration, the specific activity of the root nodules did not differ significantly between the two treatments. That observation is consistent with the data in Table III and contrasts with the response reported for CO₂-enrichment treatments of older soybeans (3-5).

The present experiments indicate that, under constant environmental conditions, soybean root nodule activity was not limited by whole-plant photosynthate production. Instead, the soybean plants under the normal light/dark cycles and CO₂ concentrations maintained a daily balance between substrate production and utilization similar to that reported by McCree and Kresovich (7). The term, utilization, in this context refers to both growth and maintenance uses of photosynthate. Short-term changes from the normal conditions, such as changes in ambient CO₂ concentrations or extension of the dark period, altered that balance. Long-term increases in shoot CER by CO₂ enrichment stimulated the plants to readjust and to develop a new equilibrium between daily substrate production and utilization. Thus, under controlled conditions, increases in whole-plant CER did not directly affect root nodule activity but did alter the 24-h balance between substrate production and utilization, so that plant and nodule growth was stimulated and N₂ fixation on a whole plant basis increased. Data from subterranean clover grown under constant conditions (15) indicate that a similar 24-h balance may occur in that species when it is dependent on symbiotic N_2 fixation. Nodule activity in peas, however, can be more directly dependent on current photosynthetic activity (10). The concept of a 24-h balance between substrate production and utilization in soybeans grown under controlled environmental conditions is consistent with diurnal patterns of leaf carbohydrate level (17) and evidence for programmable rates of starch accumulation reported by Chatterton and Silvius (1). Some of the differences between the short-term responses of nodule activity to changes in photosynthesis reported in the present research and those reported by others (3-5, 8, 16)may be a function of the relative capacity of the plant to arrive at a balance between photosynthate production and utilization under varying field conditions, compared to controlled conditions. Similarly, correlations between short-term measurements of CER and root nodule activity (14) probably reflect a longer-term equilibrium between photosynthate production in the leaves and utilization by the rest of the plant, including the nodules.

Results from this study clearly establish three points. First, soybean root nodules on intact, growing plants can function at a maximum rate that is not limited by shoot CER. Second, root nodule acetylene-reduction activity of growth-chamber-grown plants is in equilibrium with shoot CER and other plant processes and cannot be perturbed by short-term increases in shoot CER. Third, long-term enhancement of shoot CER produces a coordinate increase in plant growth, nodule growth, and symbiotic N₂ fixation after several days. These results do not argue against such applied objectives as increasing photosynthetic efficiency or decreasing photorespiration, but they indicate that increases in N₂ fixation resulting from a greater shoot CER will be balanced by shoot and nodule growth during vegetative and early reproductive stages of development. Whether translocation or partitioning of photosynthate to root nodules is a major factor limiting symbiotic N₂ fixation is not addressed by these results and deserves further study.

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