

How do carbon dioxide and ozone affect the basal respiration and soil microbial population in the rhizosphere of *Glycine max*?

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ABSTRACT

This study was conducted as part of a climate change research project at USDA-BARC (United States Department of Agriculture-Beltsville Agriculture Research Center, College Park, MD, USA) to determine the effect of increased tropospheric CO₂ and O₃ on responses of soil basal respiration rates (BR) and microbial population (MP) in soybeans (*Glycine max*) rhizosphere soil. The plants cultivars (Essex and Forrest) were grown to a full-season in 16 open-top chambers supplied with four air quality treatments; charcoal-filtered (CF) air as a control at 350 ± 5 µL CO₂ L⁻¹, CF (350 ± 5 µL CO₂ L⁻¹) + 150 ± 5 µL CO₂ L⁻¹, non-filtered (NF) air (25 ± 5 nL O₃ L⁻¹) + 30 ± 5 nL O₃ L⁻¹, and NF + 30 ± 5 nL O₃ L⁻¹ + 150 ± 5 µL CO₂ L⁻¹ at two soil moisture regimes (irrigated and sheltered). The BR values were determined during four growth stages (pre-cultivation, flowering, early grainfill, and late grainfill) at three times during the day (morning, noon, and afternoon) for each stage. Also, specific maintenance respiration rates (qCO₂) were calculated for all growth stages. Microbial populations (MP) of rhizosphere soil were recorded once, at midday, for all growth stages. Significant changes in BR, qCO₂, and MP were observed for all treatments. The greatest increases in BR and MP, and the least in qCO₂, were found at early grainfill stage treatments. The deterrence in BR and MP values were stimulated by the increase of CO₂ singly, or in combination with high O₃ exposures compared to CF control. Reductions in BR and MP observed for the NF + 30 ± 5 nL O₃ L⁻¹ treatments during flowering and early grainfill stages, thus suggests that O₃ injury can reduce the BR by decreasing the activity of microbes in soil. This study suggests that carbon dioxide could decrease the harmful effect of ozone on soybean plants.

KEYWORDS: Carbon dioxide, ozone, respiration, microbes, soybeans.

INTRODUCTION

Plants like soybeans are in an atmosphere that supplies CO₂ for carbon fixation and the production of carbon-containing compounds to maintain the structure and function of cells; and O₂ to oxidize these carbon compounds, producing cellular energy. In recent human history, human activities have produced increases in both CO₂ and O₃, a highly reactive form of O₂. Singly, O₃ and CO₂, generally have either "detrimental" or "beneficial" effects on plants. However, the combined impact of both gases on vegetation is uncertain despite their likely co-occurrence in the future. Since the end of the 18th century, the Industrial Revolution and the associated increase in the human population and need for energy and raw materials, has resulted in increasing levels of atmospheric pollutants. Previously, the positive feedback between a growing human population and release of pollutants has had significant impact on a regional level due to highly reactive pollutants such as O₃, but now has reached a global level of importance due to large increases in other greenhouse gases, especially CO₂.

On a global basis, over the last few decades, O₃ concentrations have increased at between 1 to 2% per year (Fishman 1991) and concentrations are expected to continue to increase. Based on patterns of expected emission of precursors, Hough & Derwent (1990) suggested that concentrations would increase 20-50% between 1990 and 2020 in lower

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latitudes (36 to 42 degrees N) while at high latitudes (48-56 degrees N) concentrations would increase 10-28%. Regional levels of O₃ are likely to continue increasing in major cities (e.g. Mexico City, and likely Beijing, Bombay and Cairo) that continue to show rapid population growth and use of fossil fuels in automobiles and industry (Yunus *et al.* 1996). Furthermore, the geographic extent of the effect of O₃ on terrestrial ecosystems is likely to increase as more countries, particularly developing ones, become more industrialized or have more managed agriculture (Chameides *et al.* 1994).

The atmospheric CO₂ concentration has been increasing dramatically for over a century. Other greenhouse gases (e.g. CH₄, N₂O, O₃) are increasing in concentration at rates specific to each gas. Associated with the increases in CO₂ and other greenhouse gases, the IPCC (Intergovernmental Panel on climate change) estimate that global air temperatures will increase (Houghton *et al.* 1996). The potential increase in global temperature and the likely associated changes in precipitation amounts, type, and pattern; humidity, and possibly other climatic factors that could have significant impacts on the world's ecosystems and, consequently, human health and welfare.

Individual species do not exist in isolation, but as members of an ecosystem, i.e., all organisms and their nonliving environment in a given area (Odum 1971). To date most studies on air pollutant effects have focused on responses of individual components, almost entirely on a single species level and not in terms of ecosystem function. While this level of analysis has allowed for a "snapshot" of the status of an ecosystem at one point in time, it cannot provide for estimation of long-term responses. Thus we need to understand implications of pollutants for ecosystem health more fully.

Terrestrial ecosystems provide many goods and services for humans. Agro-ecosystems provide food and fiber whereas natural ecosystems, especially forests, provide timber, fuel, and non-timber products; protection of soils; sources of water; conservation of biodiversity; and recreation. There is much debate about how the various roles of Agro-ecosystems and natural terrestrial ecosystems in global phenomena will be affected by future changes in climate and a changed atmospheric composition (Solomon *et al.* 1996). Global vegetation models that have been used to simulate the transient response of the terrestrial biosphere to changes in climate and CO₂ have generally shown an eventual increase in C storage on the land after a substantial loss during the transient phase (Schimel *et al.* 1994). None of these studies have included effects of other atmospheric pollutants such as O₃, which may counter effects of a changed climate.

As O₃ and CO₂ increase globally, changes in the flux of C to and from soils (respiration) and cycling of C and N will occur. This has important implications for the functioning of ecosystems because soil C and nutrient cycles are closely associated. Since terrestrial soils contain about 71% of total terrestrial C stocks (Odum 1971), any change in the net flux of carbon into or out of soils may have major repercussions on atmospheric CO₂ concentrations and the potential for global change. As the most biologically active portion of soil, the rhizosphere (the soil immediately adjacent to plant roots) is likely to be affected most by environmental stresses. Therefore, it is essential to understand how specific stressors will affect the rhizosphere, which acts as an interface between primary carbon processes and primary nutrient and water processes. To study the rhizosphere, one needs to consider its biology and ecology as an integrated system. The interaction of biology, ecology, chemistry and physics within the mineral soil matrix creates the habitats found in soil (Rygiewicz & Ingham 1998). The biotic community below-ground includes bacteria, fungi, etc., which serve various functions in maintaining biological, physical and chemical characteristics of the soil, and all are dependent on energy inputs from plant residues for their maintenance (Kuikman, *et al.* 1990).

The main objective of this study is to answer the following question through focusing on the microbial respiration in rhizosphere soil: will rising global atmospheric CO₂ counteract the regional or local detrimental effects of atmospheric O₃ on soybean plants?

MATERIALS AND METHODS

Research Facility: The research, using open-top chambers, was conducted at the United States Department of Agriculture (USDA) Beltsville Agricultural Research Center (BARC), Beltsville, MD, USA. The site was located on a Codorus silt loam soil containing about 40% sand, 21% clay and 39% silt, with a PH of 6.2. The treatments were begun as soon as plants were visible. Using Diazinon 50 W at 0.7 kg/ha in solution, the soybean plants were sprayed one or more times during its growth to control weeds.

Soybeans (cv. Essex and Forrest) were planted in June 2000 in 4m x 5m plots in rows spaced 0.5 m apart with seeds spaced 10 cm apart for over 200 000 ha. The plots were covered in late June with 3 m diameter open-top chambers (Heagle *et al.* 1973) as soon as uniform plant stands were assured and sprinkler irrigation units had been installed in the center of each chamber. Atmospheric treatments consisted of charcoal-filtered (CF) air as a control at $350 \pm 5 \mu\text{L CO}_2 \text{ L}^{-1}$, CF ($350 \pm 5 \mu\text{L CO}_2 \text{ L}^{-1}$) + $150 \pm 5 \mu\text{L CO}_2 \text{ L}^{-1}$, non-filtered (NF) air ($25 \pm 5 \text{ nL O}_3 \text{ L}^{-1}$) + $30 \pm 5 \text{ nL O}_3 \text{ L}^{-1}$, and NF + $30 \pm 5 \text{ nL O}_3 \text{ L}^{-1}$ + $150 \pm 5 \mu\text{L CO}_2 \text{ L}^{-1}$ in a complete factorial design for a total of four atmospheric environments.

The treatments were arranged in a randomized complete block design and replicated twice for a total of 16 chambers (4x2x2). Half of these chambers were equipped with moveable rainfall shelters to exclude rainfall (sheltered soil moisture). Another half (irrigated plots) received natural rainfall and was irrigated to maintain moisture (wet -0.05 MPa x dry -1.5 Mpa). The CO₂ treatments were applied 12 h a day (05:00-17:00 h EST), 7 days a week and the O₃ treatments were imposed 7 h a day (09:00- 16:00 h EST), 5 days a week for 12 weeks. The CO₂ was supplied from cylinder CO₂ and the O₃ was synthesized from cylinder O₂ using a Griffin O₃ Generator (Griffin Technics Crop. Lodi, NJ). Both gases were injected into the chamber blowers immediately upstream from the fans. The flow of CO₂ to each chamber was monitored through glass flow meters and the individual rates were checked daily. The CO₂ was monitored in the stream prior to contacting the plants using a Beckman Model 315 B Infrared Analyzer. The CO₂ monitor calibrated using certified CO₂ standards in N₂ gas purchased from Air Products and Chemical Co., Washington, DC. The air stream CO₂ concentrations were measured on a biweekly basis and adjustments to flow rates were made if necessary. Chamber O₃ concentrations were monitored hourly during the treatment periods using a Thermo Electron Model 49, UV photometric O₃ Analyzer (Thermo Electron Corp., Hopkinton, MA). The O₃ meters were calibrated using a Dasibi Model 1003PC (Dasibi Environmental Corp., Glendale, CA). The chamber air samples were collected using Teflon^R tubing attached through a switching device to a central vacuum system. Samples lines in the chambers were adjusted weekly to about 0.2 m above the canopy throughout the growing season.

Soil collection: Soil samples were collected in late 1999 to a depth of 15 cm using a 1.9 cm i.d. soil probe. Six cores were randomly collected from each site of the plot or from each soil cultivars. The cores were pooled and mixed in the field immediately after a site was sampled, and placed in tightly sealed plastic bags. Soils were transported from the field site in plastic bags kept on ice in a dark cooler. The soil cores were gently sieved to pass through a 4 mm mesh to remove stones, roots and large organic residues.

Basal respiration rates (BR): The BR ($\text{M CO}_2\text{-C m}^{-3} \text{ d}^{-1}$) was measured as the average CO₂-evolution of soil after an incubation period of 10 days. The BR values were determined during four growth stages (pre-cultivation, flowering, early grainfill, and late grainfill) on three times

through the light hours of the day [morning (10am), noon (12pm), and afternoon (2pm)] for each stage. The BR rates were calculated as follow:

$$BR = (CO_2 - C_{UNFUM} - CO_2 - C_{AIR}) / 10 \text{ days}$$

Where $CO_2 - C_{UNFUM}$ is the evolution of CO_2 during 10 days incubation of non-amended soil and $CO_2 - C_{AIR}$ is the CO_2 in a blank mason jar.

The BR method of Van de Werf and Verstrate (1987) was modified as follow: About 20 g oven-dried equivalent (ODE) of 2 mm sieved non-amended homogenized soil adjusted to 60% water-filled porosity (WFP) was placed in each of two 50 mL glass beakers. The soil in one beaker was amended with a glass vial containing 10 mL of distilled water to maintain humidity and a plastic vial containing 10 mL of 1 M NaOH to trap evolved CO_2 in a 1 L mason jar. Another group of jars were used as a control without using soil samples. The jars were sealed and incubated in the dark for 10 days at $25 \pm 1^\circ C$. Following the incubation period, the Na_2CO_3 formed in the vial containing 1 M NaOH was precipitated as $Ba CO_3$ by addition of 1 M $Ba Cl_2$. the remaining NaOH in each vial was titrated to the phenolphthalein endpoint with a standardized 1 M HCl solution (Islam et al. 2000).

The specific maintenance respiration rates (qCO_2): The qCO_2 is the CO_2 release per unit of microbial biomass in soil. They were calculated as mean BR rates over total microbial biomass (BR/C_{TMB}) ($M CO_2 - C m^{-3} d^{-1} C_{TMB}^{-1}$) using method of Anderson and Gray (1991).

Microbial population (MP): Microbe's number for variable soil rhizosphere in soybean was enumerated by spiral plating assay (Spiral System Inc., Model D, Cincinnati, OH). Microbial populations (MP) of rhizosphere soil collected once time for all growth stages through mid of the day. Bacterial and fungal enumeration by the spiral plating method involve the counting of colonies in the outer region of the plate, where the colonies are well separated, and dividing these counts by the volume of sample deposited within the counting region. Agar plate medium was used to prepare the plates. The same quantity of medium was poured into all plates so that the same height of agar was presented to the spiral plate stylus tip to maintain contact angle.

Soil samples were collected from the root area after shacking firmly. Ten grams of soil were added to 90 mL of buffer and homogenized in blender at 22000 rpm for one min. and one min., for settling. Sterile buffered solution tubes, containing 9 mL, were prepared to make a series dilutions. One mL of soil dilutions amended with bacterial and fungal organisms was used to inoculate 9 mL buffered solution, creating a 1:10 dilution.

Stock A: 50.0 g of $MgSO_4 \cdot 7H_2O$ /liter. For buffered H_2O , 5.0 mL stock was used per liter. Stock B: 34.0 g of KH_2PO_4 /litre. For buffered H_2O , 1.25 mL stock was used per liter.

The platter was connected to appropriate electric and vacuum sources, and set in the automatic model. A marked 5 mL disposable cup was filled with ethanol and another cup containing sterile distilled water was placed in the post mounted cup holder. The stylus was sanitized by immersing the stylus tip three times into the alcohol with the pinch valve open (valve switch on). The stylus was then rinsed by immersing the tip into the water. The water was pulled through the system once the valve was closed. The serial dilution of each treatment was poured into a disposable cup and placed in the cup holder. The stylus was lowered into the sample and the valve was opened until there was a continuous column of liquid sample without bubbles in the sight glass (about 2-3 sec.). The valve was turned off, the stylus was raised, and the sample holder was moved out of the way. The dish was placed on the turntable, the stylus was lowered onto the medium, and pressing the start switch started the platter. When the plating cycle was completed, the stylus was raised automatically and the plate was removed and its lid was replaced.

The plates were inverted and incubated at $28^\circ C$ in the dark for 4 days. After incubation, sector was chosen and counting of colonies was begun from the outer edge of the first segment towards the center until 20 colonies have been counted. A similar area on the

opposite side of the plate was counted and the sample volume deposited in those two areas divided the colonies counted from both sides. The volumes of samples associated with each portion were given in sectors (8, 9, 10, 11, 12, 13, and complete plate) as 100 mm plate size μL deposited i.e. 1.145, 2.798, 5.186, 8.634, 13.618, 24.151, and 48.302 respectively. The volume constants found for segments, and then multiply by 1000 for CFU/mL divided the number of counted colonies from both sides of the plate.

Statistical analysis: Data were analyzed using analysis of variance (ANOVA) procedures for factorial design $4 \times 2 \times 2$. The least Significance Difference (LSD) evaluated the mean differences between the four air quality treatments. Statistical Analysis System (SAS) was performed all data statistical analysis (SAS Institute, 1990).

RESULTS

Meteorological data including temperature, wind velocity, precipitation, and radiation values during the day for collecting samples obtained from a location adjacent to the experimental site are summarized in Fig. 1.

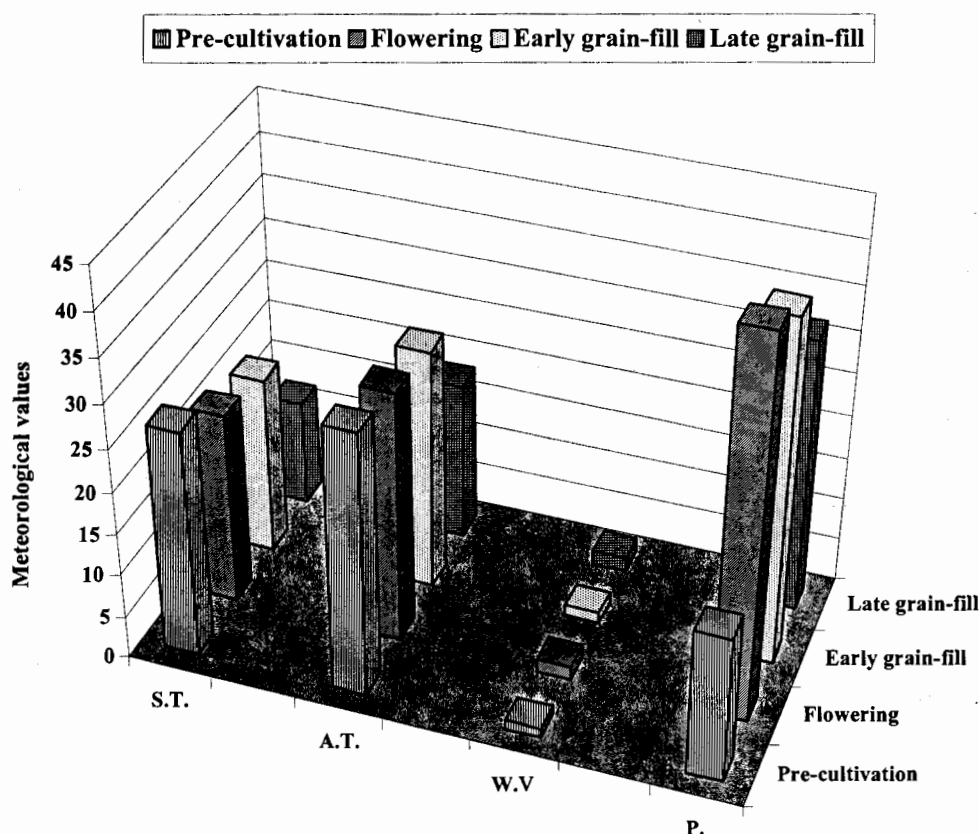


Fig. 1: Soil temperature (ST = °C), air temperature (AT = °C), wind velocity (WV = ms⁻¹) and precipitation (P = mm) values during the day for collecting samples.

The effects of atmospheric CO₂, O₃ and moisture regimes on BR rates for soil supporting soybean at four growth stages of plant development [pre-cultivation, flowering, early podfill (early grainfill), and grainfill (podfill)] are illustrated in Fig. 2. Moisture regimes produced significant effects at the pre-cultivation stage and at the early pod stage of development with lower values under dry treatments. Air quality treatments caused significant differences during the day at all four growth stages with elevated CO₂

producing higher CO₂ flux, and elevated O₃ concentrations giving lower values compared to carbon-filtered controls. Moreover, the afternoon data showed much higher levels than the early morning ones.

The effects of cultivars of soybean on respiration rates were not significant in the major cases. The Forrest cultivar exhibited slightly higher rates of respiration than samples collected from Essex. The effect of soil moisture on CO₂ flux rates in soil supporting soybean showed significant increases in wet conditions during all times of the early grainfill stage, but especially in the morning of pre-cultivation stage (Fig. 2). Generally, elevated CO₂ treatments increased the rates of flux while O₃ treatments exhibited reductions in all levels. The BR data recorded the highest values for all growth stages are the afternoon. Soils under high O₃ treatments typically showed lower levels of respiration rates than carbon-filtered controls, with only two instances where the differences were not significant, afternoon of pre-cultivation and noon of late grainfill stages. The combination of both CO₂ and O₃ at high concentrations slightly increased the levels of respiration at flowering and early grainfill stages, especially under high moisture concentrations.

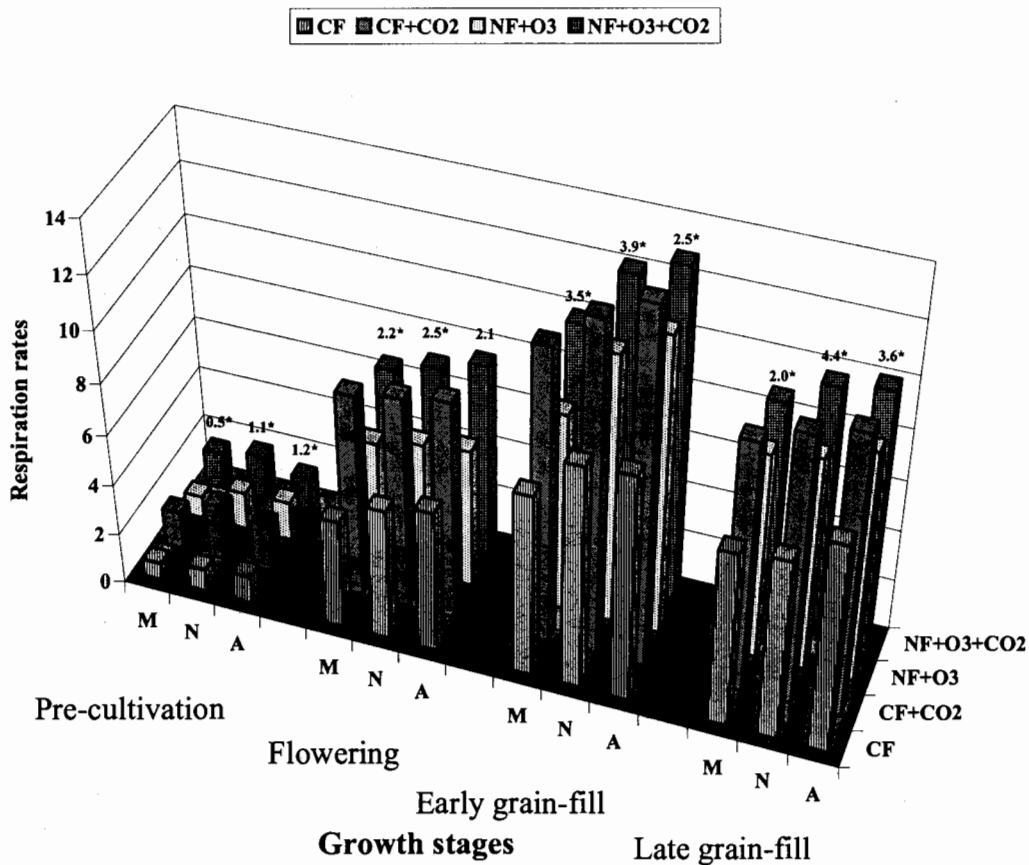


Fig. (2): Basal respiration rates values (M CO₂-C m⁻³ d⁻¹) of soil supporting soybean plants under atmospheric CO₂ and O₃ treatments. (M = morning, N = noon, A = afternoon, CF = carbon filtered, NF = non filtered air. * = LSD values).

The effects of air quality and soil moisture levels on BR rate values were largely significant under soil moisture treatments. Respiration levels in the all growth stages were lower under dry conditions during all times of the day. With regard to air quality treatments, exposure to high O₃ significantly reduced the levels of respiration while high CO₂ increased levels compared to carbon-filtered air (Fig. 2). The highest respiration rates for all growth

stages were observed during early pod development. With respect to the combination of elevated CO₂ and O₃; respiration levels at all times of the day were comparable if not slightly larger, than those observed in the carbon-filtered control treatments.

The impact of elevated CO₂, O₃ and soil moisture regimes on qCO₂ soils under soybean roots are summarized in Fig. 3. In term of air quality treatment effects, the results were typically significant for CO₂, showing decreases in all growth stages and at all times during the day. The effects of O₃ treatments were generally significant compared to the carbon-filtered air controls but wise insignificant in the mornings of flowering and early podfill stages. The effects of combined high CO₂ and high O₃ included a decrease in qCO₂ levels for all growth stages and especially in the flowering and early podfill stages compared to O₃ treatments.

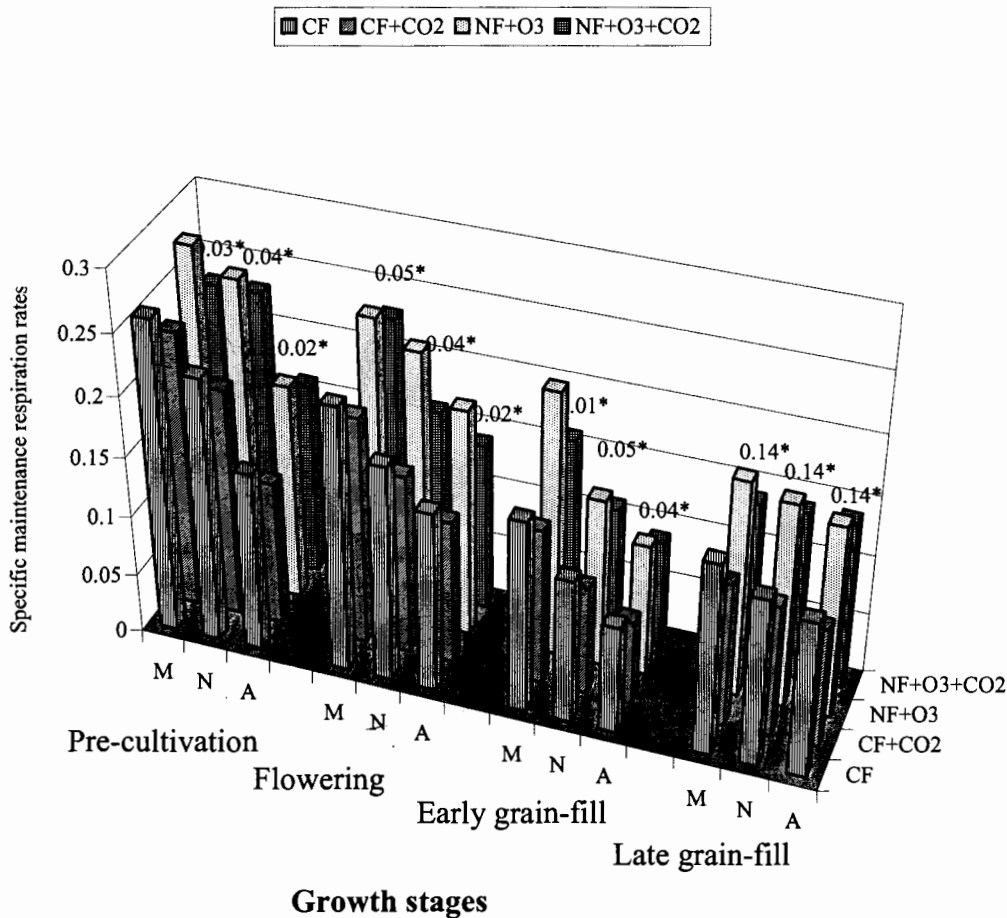


Fig. 3: Specific maintenance respiration rates values (qCO₂) of soil supporting soybean plants under atmospheric CO₂ and O₃ treatments. (M = morning, N = noon, A = afternoon, CF = carbon filtered, NF = non filtered air. * = LSD values).

The soybean cultivar results indicate significant differences for all growth stages except the pre-cultivation stage. The moisture treatments exhibited minimal significant difference between the two regimes. In terms of air quality treatment effects and moisture, high CO₂ concentrations caused decreases in qCO₂ in both pre-cultivation and flowering stages as compared to carbon-filtered air controls. Elevated O₃ treatments showed a significant increase of qCO₂ (Fig. 3). The combination of elevated CO₂ and O₃ produced

significantly higher levels of qCO_2 compared to high CO_2 treatments. There were no observed interaction effects between the four air quality treatments and soil moisture regimes with similar patterns of responses being found under both moisture treatments for early podfill and late podfill stages.

The MP counts in the soybean rhizosphere soils from the open-top chamber under atmospheric CO_2 or O_3 enrichments and two moisture regimes over afternoon periods are shown in Fig. 4. Fungal populations were not significantly affected by the moisture treatments while bacterial results were significantly different. Atmospheric CO_2 treatments typically increased the MPs in the soil while ozone enrichment tended to inhibit the growth of both types of microorganisms in the soil rhizosphere. The combination treatment of CF + CO_2 and high O_3 treatments showed trends for increased populations, with few samples appearing significantly higher than the charcoal filtered air treatments.

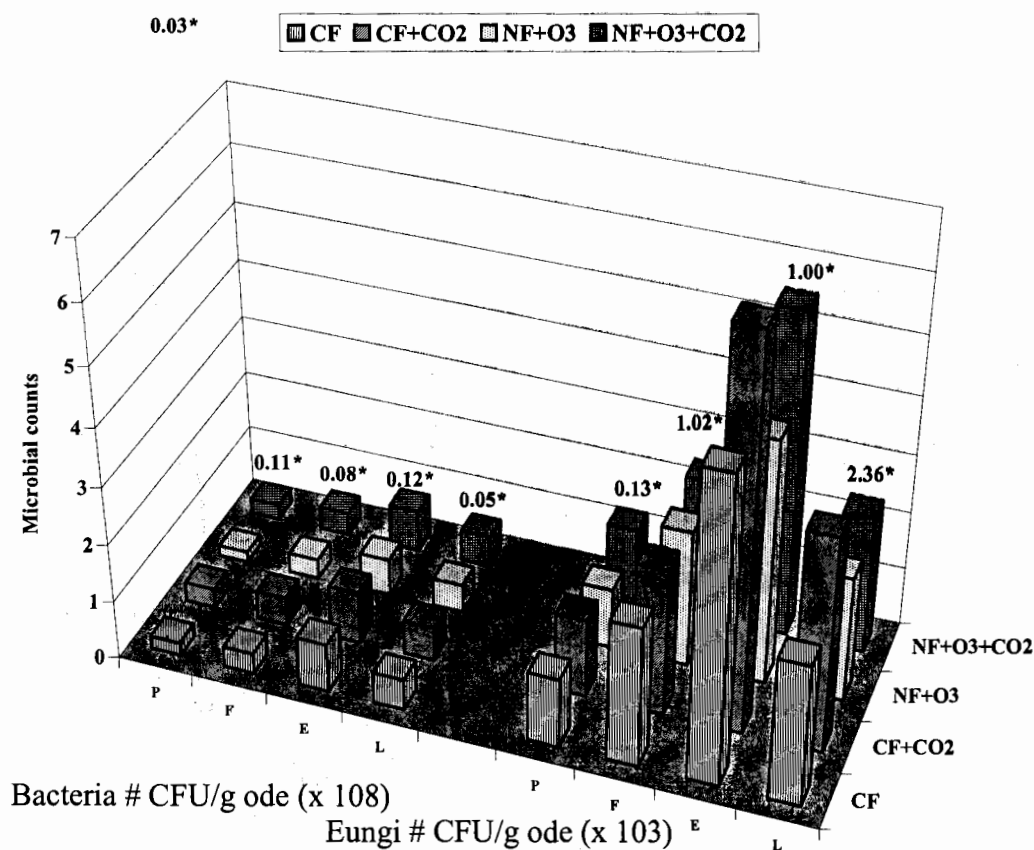


Fig. 4: Microbial counts values of soil supporting soybean plants under atmospheric CO_2 and O_3 treatments. (P = pre-flowering, F = flowering, E = early grain-fill, L = late grain-fill CFU = colony forming unit, Ode = oven dry equivalent, CF = carbon filtered, NF = non filtered air. * = LSD values).

The combination of air quality treatments with moisture regimes generally showed increased MPs under CF + CO_2 for both moisture conditions; however, NF + O_3 treatment results were consistently lower than the charcoal filtered air controls although were normally not significantly different. Although the data from both moisture regimes were somewhat varied over the periods of growth, the patterns of results appeared generally similar under the two moisture regimes. The interaction of air quality treatments vs. moisture treatments was

significant in most cases for bacterial counts and fungal numbers in soils (Fig. 4). Under the wet treatments, fungal counts for the elevated CO₂ treatments were significantly higher than CF controls; however, results for high O₃ treatments were all comparable to CF controls, except for a few results during the late grain fill stages. Additionally, the combination of high O₃ and CO₂ treatments had fungal counts larger than CF controls under the dry treatments. The high concentration of O₃ and CO₂ treatments stimulated the fungal counts compared to CF controls in the combined results, while the results for the NF + O₃ + CO₂ treatments were generally insignificant. Bacterial counts were also stimulated by the elevated CO₂ treatments but high O₃ and NF + O₃ + CO₂ treatment effects were largely non significant in pre-cultivation and late pod fill stages under dry conditions.

DISCUSSION

The present study investigates the dependence of BR, MP, and qCO₂ on climatic conditions. The data showed that strong responses to BR and MP under the four air quality treatments gradually from pre-cultivation to early grain-fill stages and decreased through the late grainfill stage. The treatments of tropospheric O₃ decreased CO₂ evolution in wet moisture regimes. Significant interactions of CO₂ x O₃ enrichments with moisture were observed for qCO₂ during all growth stages. The results for MP exhibited similar patterns to that for BR, where significant increases were found in both bacteria and fungi in rhizosphere soil subjected to elevated CO₂ effects and effective decreases under high O₃ concentrations. These results agree with those obtained by Insam (1990); Edward (1991); Koizumi *et al.* (1991); Tingey *et al.* (1995); Cheng *et al.* (1996); Schortemeyer *et al.* (1997); North & Nobel (1997).

Generally, significant relationships were found between the effects of CO₂ and O₃ treatments, and CO₂ fluxes and microbial numbers. The functional relationship between BR and MP is not fully understood. Sparling (1991) considered BR to be representative of the active part of MP. Anderson and Domsch (1985), however, viewed the BR as reflecting the activity of whole microbial activity. Typically, a high qCO₂ is found in soils with a recent input of easily degradable substrate. Such substrates would induce a microflora that usually respired more CO₂ per unit degradable C (Islam *et al.*, 2000). The relationship between BR and qCO₂ were found to be linked to climatic conditions. Part of the climatic effect may be explained by an altered quantity of metabolizable substrates due to an influence on primary production or substrate allocation to the roots and decomposition as such in response to climatic conditions. Both factors affected the mediator of decomposition, MP; BR. Significant relationships between qCO₂ and the stimulation of microbial activity may be attributed to the translocation of photosynthetic compounds under ground. The suitable time for detecting BR and MP is the period after 12pm because higher temperatures correlate with soil surface CO₂ fluxes and accelerate the development of a soil (Jensen *et al.* 1996).

Soil surface CO₂ evolutions can originate from any ecosystem carbon. The majority of CO₂ fluxes that reach plants could originate from the soil and be related to daily net photosynthetic rates (Monteith *et al.* 1964). The respiration be produced in soil by the activity of roots, and microbes and affected by soil and air temperatures, soil moisture content, wind velocity, is precipitation and solar radiation (Rochette *et al.* 1997). Also, Kassim *et al.* (1982) suggested that lower metabolic efficiency is mean high maintenance respiration. Basal soil respiration values are inversely proportional to qCO₂ due to efficient assimilation of organic C by higher proportions to activity of microbes (Islam *et al.* 2000).

Plant and soil-associated biota such as bacteria, fungi, etc. play important roles in nutrient cycling and plant productivity in natural and managed ecosystems. Penetration of O₃ into soil supporting soybean plants is believed to be limited essentially to the soil surface.

In *Acer saccharum*, a decrease in the number of arbuscules, an increase in the number of vesicles in roots and decreases in the numbers of rhizosphere bacteria and non-arbuscular mycorrhizal fungi in soil were associated with increased levels of tropospheric O₃, increased UV-B radiation and increased levels of flavinoids, tannins and lignins (Rozema *et al.* 1997). Edwards (1991) concluded that decreased respiration rates in the soil surrounding roots of the O₃-exposed plants imply that root-derived organic materials available for microbial proliferation may have reduced as an indirect result of the O₃-exposure.

Increases in global atmospheric CO₂ will not only directly affect the growth of plants, but might also alter the living conditions for soil biota. Elevated CO₂ has been reported to affect on population sizes of bacteria in the rhizosphere widely. Rogers *et al.* (1994) include reports of increases in total bacterial counts in the rhizosphere of *Gossypium hirsutum* exposed in Free Air CO₂ Enriched (FACE) rings. Increases in microbial biomass in the rhizosphere soil surrounding *Populus grandidentata* roots of plants exposed in open top chambers to CO₂ treatments have also reported (Zak *et al.* 1996). Norby (1987) reported an increase in arbuscular mycorrhizal (AM) fungi infection in *Trifolium*, which was attributed to increased root mass and AM fungal entry points, rather than to changes in root exudate levels or composition. Generally, Schortemeyer *et al.* (1997) indicated that in legume crops, at least in terms of inoculum quality in the rhizosphere soil, symbiotic nitrogen-fixing organisms might be favored by elevated atmospheric CO₂ concentrations.

The effects of CO₂ and O₃ interactions on soil biota are few. Perez-Soba *et al.* (1995) studied the effects of joint CO₂ and O₃ exposures on mycorrhizal infection of Scots pine saplings. High O₃ reduced mycorrhizal infection of roots; however, there was no CO₂ and O₃ interaction, as elevated CO₂ had no effect on mycorrhizal infection at either O₃ level.

Given the broad range of responses both within and between plant genera and species, differences in sampling methodology, differences in the exposure regimes and length of studies, it is clear that more research is needed to clarify the nature and mechanism of the effects of O₃ and CO₂ on plant and soil associated microbiota. Assessment of the effects of O₃ and CO₂ on non-symbiotic N₂ fixation, on non-symbiotic, pathogenic and saprophytic fungi, and on soil invertebrates and on the quality, quantity and rates of decomposition of litter are urgently needed. Collectively, availability of these diverse types of data will enhance our ability to identify and develop measures of ecosystem status and function. The data will also be useful to develop and validate predictive models of the effects of elevated CO₂ and O₃ or other stressors on ecosystem integrity. With increased industrialization and concomitant increases in levels of tropospheric O₃, information on O₃ effects on plant and soil associated biota is needed to help identify potential indicators of O₃ damage. The information may also be useful to understand the basic mechanisms by which the damage occurs and to develop strategies to minimize or mitigate adverse effects of ozone exposure in natural and managed ecosystems. Effects of ozone on plant-associated microbes in rhizosphere soil indicate sometimes-contrasting results, and ones, which may be genus or even plant species specific.

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الملخص العربي

هل يؤثر غاز ثاني أكسيد الكربون والأوزون على تنفس التربة القاعدي ومحتواها من الميكروبات في الجزء السطحي من تربة نبات فول الصويا

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تهدف الدراسة التي اقيمت كجزء من مشروع بحث التغيرات المناخية في الولايات المتحدة الأمريكية بولاية ميريلاند تحت اشراف وزارة الزراعة الي تحديد مدي تاثير الزيادة في غاز ثاني اكسيد الكربون و الاوزون علي استجابة معدل تنفس التربة القاعدي ومحتواها من الميكروبات في الجزء السطحي من تربة نبات فول الصويا. زرعت هذه النباتات لموسم كامل في ١٦ صوبه نباتية مفتوحة تحت تاثير اربعة انواع من المعاملات الهوائية وهي: هواء مرشح عند تركيز 350 ± 50 ميكروليتر غاز ثاني اكسيد الكربون، هواء مرشح مضاف اليه 150 ± 50 ميكروليتر غاز ثاني اكسيد الكربون، هواء غير مرشح مضاف اليه 30 ± 50 نانو ليتر غاز الاوزون، هواء غير مرشح مضاف اليه 30 ± 50 نانو ليتر غاز الاوزون و مضاف اليه 150 ± 50 ميكروليتر غاز ثاني اكسيد الكربون تحت تاثير اثنين من ظروف رطوبة التربة وهما جيدة ومحدودة الرطوبه. ولقد تم تعيين معدلات التنفس القاعدي اثناء اربعة من مراحل النمو وهي: مرحلة قبل الزراعة، مرحلة الازهار، بداية مرحلة امتلاء القرون، نهاية مرحلة امتلاء القرون علي ثلاث فترات اثناء اليوم: في الصباح، اثناء الظهر، بعد الظهر لكل مرحلة من مراحل النمو. وقد تم ايضا حساب معدلات صيانة التنفس المتخصصة لكل مرحلة من مراحل النمو. جمعت العينات الخاصة بالتجمع الميكروبي في الجزء العلوي من التربة مرة واحدة اثناء منتصف اليوم لكل مرحلة من مراحل النمو. اوضحت النتائج فاعلية التغيير في معدلات التنفس القاعدي و معدلات صيانة التنفس المتخصصة ومحتواها من الميكروبات لكل مرحلة من مراحل النمو. ولقد وجد ان اعلي زيادة في معدل تنفس التربة القاعدي ومحتواها من الميكروبات في بداية مرحلة امتلاء القرون تحت التاثير العالي من غاز ثاني اكسيد الكربون بمفرده او في حالة اضافته الي التركيز العالي من الاوزون مقارنة بتركيز هواء مرشح عند تركيز 350 ± 50 ميكروليتر غاز ثاني اكسيد الكربون والعكس في حالة معدلات صيانة التنفس المتخصصة. ابرزت الدراسة ان النقص في معدل تنفس التربة القاعدي ومحتواها من الميكروبات تحت تاثير هواء غير مرشح مضاف اليه 30 ± 50 نانو ليتر غاز الاوزون اثناء مرحلة الازهار، بداية مرحلة امتلاء القرون من شأنه التاكيد علي ان التاثير المنمر للاوزون علي معدل تنفس التربة القاعدي عن طريق تثبيط النشاط الميكروبي في التربة. واثبتت الدراسة انه من الممكن ايقاف التاثير الضار للاوزون علي نبات فول الصويا باضافة تركيز عالي من ثاني اكسيد الكربون.