**Funneliformis mosseae** Improves Growth and Nutrient Accumulation in Wheat by Facilitating Soil Nutrient Uptake under Elevated CO2 at Daytime, Not Nighttime

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Abstract: The concurrent effect of elevated CO2 (eCO2) concentrations and arbuscular mycorrhizal fungi (AMF) on plant growth, carbon (C), nitrogen (N), phosphorus (P) and potassium (K) accumulations in plant and soil is largely unknown. To understand the mechanisms of eCO2 and mycorrhization on wheat (*Triticum aestivum*) performance and soil fertility, wheat seedlings were grown under four different CO2 environments for 12 weeks, including (1) ambient CO2 (ACO2, 410/460 ppm, daytime/nighttime), (2) sole daytime eCO2 (DeCO2, 550/460 ppm), (3) sole nighttime eCO2 (NeCO2, 410/610 ppm), and (4) dual or continuous daytime/nighttime eCO2 ((D + N)eCO2, 550/610 ppm), and with or without AMF (*Funneliformis mosseae*) colonization. DeCO2, NeCO2 and (D + N)eCO2 generally significantly increased shoot and root biomass, plant C, N, P and K accumulation, soil invertase and urease activity, but decreased shoot and root N, P and K concentrations, and soil available N, P and K. Compared with non-AMF, AMF effects on above-mentioned characteristics were significantly positive under ACO2, DeCO2 and (D + N)eCO2, but negative on plant biomass, C, N, P and K accumulation under NeCO2. Overall, AMF colonization alleviated soil nutrient constraints on plant responses to DeCO2, while NeCO2 decreased AMF’s beneficial effects on plants. These results demonstrated that an integration of AMF’s benefits to plants under factual field DeCO2 and/or NeCO2 will be critical for managing the long-term consequence of future CO2 rising on global cropping systems.

Keywords: arbuscular mycorrhiza; biomass production; enzyme activity; nitrogen; phosphorus; potassium

1. Introduction

An increase in atmospheric carbon dioxide (ACO2) concentration is one of the most important environmental factors reflecting global climate change [1]. The ACO2 concentration has been increased from 280 ppm during the industrial revolution to 419.64 ppm (https://www.co2.earth Accessed on 5 June 2021), and could reach ~550 ppm in the next 50 years [1]. Elevated CO2 (eCO2) directly influences soil–plant systems via improving plant growth [2–4]. Elevated CO2 generally exhibits ‘fertilization effects’ due to stimulation of photosynthesis and biomass accumulation in various C3 crops, including wheat (*Triticum aestivum* L.) [5–7]. Increased productivity of crops needs a large supply of nutrients (especially nitrogen, N; phosphorous, P; and potassium, K; etc.) to match their increased carbon
(C) assimilation under eCO$_2$ [8–11]. Soil nutrient availability was indeed decreased over a long-term eCO$_2$ owing to an increased nutrient demand by eCO$_2$-stimulated growth [12]. The eCO$_2$ effects were often gradually diminished when plants grew under nutrient-limited soils [13,14]. Furthermore, the beneficial effect of eCO$_2$ on biomass production can lead to changes in C, N, P, K, and other nutrient content in both plants and soils [15–18]. These changes affected agricultural ecosystem processes, including nutrient cycling [8], soil organic matter decomposition [19], and microbial processes [20,21]. Soil enzymes, which integrate information from soil microbial and biochemical statuses, could serve as indicators of various changes in the plant–soil system [22]. Thus, soil enzyme activities are also likely to change under eCO$_2$ [3,23]. It would be worthy to understand the response of aboveground biomass and the feedback of belowground components (especially soil nutrients, soil microorganisms and enzyme activities) to eCO$_2$.

Arbuscular mycorrhizal fungi (AMF) can form mutualistic associations with the roots of >80% of vascular plant species and improve plant nutrient uptake [24]. AMF are important in defining plant responses to eCO$_2$ [4,25–28]. Elevated CO$_2$ at daytime or continuously from daytime to nighttime has a positive effect on C$_3$ crops’ photosynthetic C assimilation, which increases the belowground transportation of photosynthates to support AMF symbiosis [29,30]. In turn, AMF help to alleviate the increased plant nutrient limitation associated with increased photosynthetic rates under daytime or continuous eCO$_2$ [27,30,31]. The costs and benefits to plants and AMF are a function of the balance between the C cost of fungi and nutrient supply to the plant. On the one hand, eCO$_2$ usually decreases N, P, and K concentrations in plant tissues [15,17,32], such as a decrease of 27% for N, 34% for P, and 20% for K in wheat grains [33], and 29% for N in wheat flag leaves [34]. On the other hand, eCO$_2$ improves plant N, P, and K acquisition when plants associate with AMF [4,26,35,36], which help to alleviate N-, P-, or K-limitation [10]. However, the effects of mycorrhizal association are not always positive under eCO$_2$ when the C costs of AMF outweigh their benefits to plant nutrient uptake [37]. For instance, N availability to flag leaves of the *Rhizophagus intraradices* colonized durum wheat was lower under 700 ppm eCO$_2$ than under 400 ppm CO$_2$ [34]. The mechanisms for AMF on plant performance under daytime eCO$_2$ thus remain largely unknown, let alone under nighttime eCO$_2$.

In fact, the atmospheric CO$_2$ concentration at plant height is usually higher during nighttime than during daytime [38], considering the corresponding CO$_2$ variation or fluctuation with plant photosynthesis and soil respiration, particularly in agricultural fields. The average atmospheric CO$_2$ in Australia, Japan and the USA varied from 390 ppm during daytime to 465 ppm during nighttime [38]. Indeed, a three-year (2017–2019) period of our field observation recorded that the average daily atmospheric CO$_2$ ranged from 417 ± 16 ppm at daytime and 463 ± 27 ppm at nighttime at the National Monitoring Base for Purple Soil Fertility and Fertilizer Efficiency close to the campus of Southwest University, Chongqing, China (see Figure S1 from October 2017 to March 2018). As a consequence, plants should differentially respond to such contrasting daytime or nighttime atmospheric CO$_2$ concentrations [39]. Thus variations in daytime and/or nighttime atmospheric CO$_2$ concentrations shall provide a closer simulation of currently atmospheric CO$_2$ conditions that plants will respond to in the near future. However, owing to the likely extra cost and maintenance of CO$_2$ gas supply, only a few studies have examined the different responses of plant growth and yield between contrasting daytime and nighttime eCO$_2$ concentrations [40–44]. For instance, it was daytime, not nighttime eCO$_2$, that improved *Morus alba* growth [39]. The biomass production in *Amaranthus retroflexus* and *Zea mays* was significantly reduced under 700 ppm nighttime eCO$_2$ than under 370 ppm nighttime eCO$_2$ [45], although such nighttime eCO$_2$ effects on plant growth were plant species-specific [46]. The contrasting positive or negative effects of nighttime eCO$_2$ on plant growth would result from dark respiration [47,48]. It is suggested that dark respiration was decreased by 20–45% under 700–1000 ppm eCO$_2$ at nighttime in the short-term [41,43,49]. In contrast, dark respiration usually, but not always, increased in the long-term such CO$_2$...
Mechanisms of eCO₂ at nighttime that may affect plant growth and C assimilation have not been established yet. Questions hence arise as to whether the plant nutrient demand could be increased under eCO₂ at nighttime, and whether AMF symbiosis could play the same role in C and nutrient balance under eCO₂ at daytime or nighttime. Such information is essential for understanding the mechanisms affecting C and N dynamics under future CO₂-increasing scenarios.

We therefore designed an environment-controlled system, which has minimal impact on light, air temperature and humidity, while providing either on-site ambient or elevated CO₂ concentrations for growing plants during daytime and/or nighttime. The objectives of the present study were to address: (1) How auto-controlled field daytime and/or nighttime eCO₂ could affect plant biomass production, and C, N, and P uptake or accumulation; (2) whether AMF effects on soil nutrient uptake could alleviate nutrient constraints on responses to eCO₂ at both daytime and nighttime; and (3) whether the interactive effects of eCO₂ and AMF on plant performance and related soil properties could be differentiated between daytime and nighttime eCO₂. In doing so, winter wheat (T. aestivum cv. Yunmai) inoculated with or without AMF was grown in soil (Eutric Regosol, FAO Soil classification system) filled pots inside environmentally controlled glass-made chambers, which had similar growth conditions except different CO₂ concentrations at daytime and/or nighttime. Plant performance and soil properties were then compared 12 weeks after sowing.

2. Materials and Methods
2.1. Experiment Design and Treatments

In a completed random arrangement, the experiment was a split plot design with atmospheric CO₂ concentrations as the main factor and mycorrhizal inoculation as the subfactor, and involved two AMF treatments: inoculated with Funneliformis mosseae and autoclaved F. mosseae (non-AMF) and four atmospheric CO₂ concentration treatments (Figure 1). Based on the on-site daily observations of 417 ± 16/463 ± 27 ppm (daytime/nighttime) between October 2017 and March 2018 (see Figure S1A,B) and an estimated ~550 ppm in the next 50 years [1], four different day and/or night CO₂ concentrations (±30 ppm) or treatments were applied: (1) ambient CO₂ (ACO₂, 410 ppm daytime/460 ppm nighttime), (2) daytime eCO₂ only (DeCO₂, 550/460 ppm), (3) nighttime eCO₂ only (NeCO₂, 410/610 ppm), and (4) continuous daytime and nighttime eCO₂ [(D + N)eCO₂, 550/610 ppm]. The respective daytime and nighttime eCO₂ concentrations were thus increased by ~33.33% of the ACO₂ treatment. Daytime was from 07:00 a.m. to 19:00 p.m. and nighttime was from 19:00 p.m. to 07:00 a.m. Each CO₂ treatment owned three chambers for a total of 12 chambers to the four CO₂ treatments in a completely randomized experimental arrangement within three blocks (Figure S2A). The arrangement of chambers was crisscrossed with 4 m apart from each other to avoid the sunshade of chambers.

![Figure 1.](image-url) A schematic diagram showing the experiment designs. NM: non-AMF; AM: inoculated AMF. (A): ACO₂, ambient CO₂; (B): DeCO₂, elevated CO₂ concentrations at daytime; (C): NeCO₂, elevated CO₂ concentrations at nighttime; and (D): (D + N)eCO₂, elevated CO₂ concentrations at both daytime and nighttime.
2.2. Experimental Facility

This study was conducted from 9 November 2017 and 2 February 2018 in a CO₂ exposure facility at the National Monitoring Base for Purple Soil Fertility and Fertilizer Efficiency (29°48′N, 106°24′E, 266.3 m above sea level) on the campus of Southwest University, Chongqing, China. The CO₂ auto-controlling facility (DSS-QZD, Qingdao Shengsen Institute of CNC Technology, Shandong, China) consists of a control system and 12 environmentally controlled chambers (Figure S2A), and the CO₂ is supplied by CO₂ cylinders (Figure S2F). The CO₂ cylinders with electric point pressure meters are connected to a CO₂ control system to maintain CO₂ gas flow into each chamber with the targeted CO₂ concentration (Figure S2E).

Each growth chamber had a rectangular floor base, supported by a steel frame hanging 50 cm above the cement ground base (Figure S2A). The bottom floors of the growth chamber are made up of polyvinyl chloride plates, and the four-sided walls and top roofs of the chamber are constructed by tempered glass (10 mm thickness, 90% light transmission rate, Yutao Glass Company, Jiulongpo, Chongqing, China) (Figure S2A). The growth chamber has a size of 1.5 m × 1.0 m × 2.5 m (length × width × height) in order to grow maize, Sorghum bicolor, Glycine max, wheat, and so forth. The electron sensors for monitoring humidity, temperature, light intensity, and CO₂ concentrations are mounted on the outer and inner surfaces of the glass wall in each chamber to monitor their variations (Figure S2B–D). The air humidity, temperature, and CO₂ concentrations are automatically controlled by their respective electronic bits and pieces (Figure S2G). The monitor’s signals are fed into proportionally integrated differential controllers that regulate the opening time within a 10 s cycle (Figure S2B,C,G). This automatic electronic controlling system can automatically regulate and instantly visualize the fluctuation of ±30 ppm CO₂ concentration, ±0.5 °C air temperature and ±5% humidity inside and outside the chamber (Figure 2 and Figure S2B,C). The targeted CO₂ concentration inside is maintained by injecting 99.99% CO₂ from the cylinder (Figure S2E,F) using a solenoid valve controlled by a mini-computer (Figure S2G). When the CO₂ concentration inside a chamber exceeds the targeted concentration, the inside air is pumped out using a pump controlled by the mini-computer and filtered with 1.0 M NaOH solution. When the humidity inside a chamber is higher than that of outside the air humidity, the inside air is pumped out using another pump controlled by the mini-computer and filtered with solid anhydrous calcium chloride. The temperature is automatically maintained at 0.5 °C variation between inside and outside the chamber using an air conditioner (Gree, Zhuhai Gree Corp., Zhuhai, China) controlled by the mini-computer.

2.3. Mycorrhizal Inoculum, Growth Soil and Plant Growth Conditions

The inoculum of AMF (Funneliformis mosseae) was purchased from the Bank of Gnome at the Institute of Plant Nutrition and Resources, Beijing Academy of Agriculture and Forestry, Beijing, China. The inoculum was a mixture of soil (50 spores per gram dry soil), mycorrhizal mycelia and root segments. The growth soil (Eutric Regosol, FAO Soil Classification System, developed from Jurassic purple shale and sandstone) was air-dried, sieved by passing through a 2 mm mesh and sterilized at 121 °C for 120 min. The pots (height/diameter = 21/17 cm) were then filled with 3.4 kg of sterilized soil. The soil (pH 6.8) had 10.56 g of organic carbon kg⁻¹, 0.66 g total N kg⁻¹, 0.61 g total P kg⁻¹, 97 mg available N kg⁻¹, 17 mg available P kg⁻¹ and 197 mg available K kg⁻¹.

Seeds of winter wheat (T. aestivum cv. Yunmai) were surface-sterilized with 10% H₂O₂ for 20 min, thoroughly rinsed with sterile water, and then pre-germinated on sterilized moist filter paper at 25/20 °C (day/night) for 36 h. Eight germinating wheat seeds were sown in one plastic pot. A total of 20 g of F. mosseae inocula were put at a 5 cm soil surface depth inside each pot, while an equal amount of autoclaved (121 °C, 0.1 Mpa, 120 min) inoculum was supplied to the non-mycorrhizal pots. A volume of 5.0 mL filtrate (0.45 µm syringe filter, Millipore Corporation, Billerica, MA, USA) from the F. mosseae inoculum was added to each non-mycorrhizal pot to minimize differences in other microbial communities.
Then, two mycorrhizal pots and two non-mycorrhizal pots were placed into each growth chamber, and thus, the three replicated chambers had a total of six replicated pots for each CO₂ concentration treatment. Except the CO₂ concentration, the chambers had similar other growth conditions, such as light, air temperature and humidity, as monitored by the above-mentioned auto-controlling facility (Figure 2). To minimize differences in growth conditions, the positions of growth pots in each chamber were rotated once a week, and shifted to another replicate chamber once fortnightly. In addition, all the pots with plants were watered once with Hoagland solution to a total of 100 mg N, 50 mg P and 75 mg K per pot and the soil moisture during the whole growth period was maintained at 70% water-holding capacity with sterilized water by routines weighing of pots once every two days.

![Figure 2. Mean temperature (A), relative humidity (B), photosynthetic active radiation (PAR, (C)) and CO₂ concentration (D,E) over the experimental period in the growth chambers. ACO₂: ambient CO₂, DeCO₂: elevated CO₂ concentrations at daytime, NeCO₂: elevated CO₂ concentrations at nighttime, (D + N)eCO₂: elevated CO₂ concentrations at both daytime and nighttime.](image)

2.4. Harvest, Sampling and Analyses

Plant and soil samples were harvested 12 weeks after sowing during the jointing stage and were combined from the two pots in each chamber as a composite sample. Plant tissues were divided into shoots (leaves and stems) and roots. Plant fresh roots were carefully washed with tap water and rinsed with deionized water. A portion of fresh roots was stored in 50% ethanol to determine root AMF colonization. The remaining fresh roots and shoots were dried at 105 °C for 30 min and then at 75 °C for >48 h until they reached a consistent dry weight. Soil samples, collected from soils that had been well-mixed from each growth pot, were divided into two parts after the removal of debris and fine roots. The first part of the soil was air-dried for >48 h for the determination of chemical properties, and the second part was immediately transferred to the laboratory and stored at −20 °C for the determination of enzyme activities.
2.5. Determination of AMF Colonization

The percentage of root AMF colonization was measured according to Brundrett et al. [51]. The roots were cut into 1.0 cm segments and cleared with 10% (w/v) KOH in a water bath at 90 °C for 20 min, rinsed in water. The cleared root segments were acidified in 0.2 M HCl for 3 min and then stained with 0.05% trypan blue. The stained segments were mounted on glass slides, and a total of 50 randomly selected root segments from each replicate were examined under a microscope. The ratio of the number of root segments that showed a fungal structure (spores, hyphae, arbuscules or vesicles) and the total number of root segments was calculated as the percentage of root AMF colonization.

2.6. Determination of C, N, P and K in Plants and Soils

The oven-dried shoot and root and air-dried soil samples were ground to fine powder for soil and plant C, N, P and K analyses. Plant C concentration was determined using the potassium dichromate–sulfuric acid oxidation method [52]. After digestion with 98% sulfuric acid and 30% hydrogen peroxide, plant N, P, and K concentrations were determined with the Kjeldahl method, the vanadium molybdate yellow colorimetric method, and flame photometry, respectively [52]. Soil available N (AN) was measured by the micro-diffusion technique after alkaline hydrolysis [52]. Soil available P (AP) was extracted with 0.5 M NaHCO_3 and then measured by the Mo-Sb anti spectrophotometric method [52]. Soil available K (AK) was extracted with 1.0 M ammonium acetate and then determined by flame photometry [52].

2.7. Determination of Soil Enzyme Activity

Soil invertase activity (mg glucose g−1 soil h−1) was determined firstly by incubating five grams of fresh soil with 1 mL toluene, 15 mL 8% (w/v) sucrose, and 5 mL phosphate buffer (pH 5.5) for 24 h at 37 °C. After incubation, 1 mL filtrate with 3 mL 3,5-dinitrosalicylic acid were then incubated in boiling water for 5 min. Subsequently, the reaction solution was diluted to 50 mL with distilled water and spectrophotometrically measured at 508 nm [53].

Soil urease activity (mg NH_4^+–N g−1 soil h−1) was determined firstly by incubating five grams of fresh soil with 1 mL toluene, 10 mL 10% urea solution (w/v), and 20 mL citrate buffer (pH 6.7) for 24 h at 37 °C. After incubation, 2 mL filtrates were mixed with 4 mL sodium phenol solution and 3 mL 0.9% (w/v) sodium hypochlorite solution in a 50 mL volumetric flask. After 20 min, the reaction solution was then diluted to 50 mL with distilled water and spectrophotometrically measured at 578 nm [53].

Neutral phosphatase activity (mg phenol g−1 h−1) was determined firstly by incubating five grams of fresh soil, 1 mL toluene and 5 mL disodium phenyl phosphate solution and 5 mL citrate buffer (pH 7.0) for 24 h at 37 °C. After incubation, 1 mL filtrate with 5 mL borate buffer (pH 9.0), 3 mL 2.5% potassium ferrocyanide (w/v) and 3 mL 0.5% 4-aminoantipyrine (w/v) were then thoroughly mixed in a 50 mL volumetric flask. Subsequently, the reaction solution was diluted to 50 mL with distilled water and spectrophotometrically measured at 570 nm [54].

2.8. Statistical Analysis

Statistical analysis was performed using SPSS 19.0 software (SPSS Inc., Chicago, IL, USA). Data were shown as mean ± standard error (SE). All response variable data (except for root colonization) were analyzed by two-factor analyses of variance (ANOVA). The factors in the two-way ANOVA were CO_2 level and arbuscular mycorrhiza. Significant differences among treatments were compared by Tukey’s Multiple Range Test at p < 0.05 using SPSS 19.0 software. Graphs were plotted using OriginPro2018 software (OriginLab Corp., Northampton, MA, USA).

3. Results

3.1. Mycorrhizal Colonization

A significantly greater percentage of root AMF colonization among CO_2 treatments (p < 0.05) ranked in AMF plants as (D + N)eCO_2 (53.60 ± 2.25) > DeCO_2 (48.31 ± 3.04) ≈
NeCO₂ (47.07 ± 3.42) > ACO₂ (40.67 ± 2.59), whereas no AMF colonization was detected in non-AMF plants.

3.2. Effects of AMF and CO₂ on Plant C, N, P and K Concentration

C concentrations in shoots were significantly higher under eCO₂ than under ACO₂ in non-AMF 12-week-old wheat seedlings, but no significant differences among DeCO₂, NeCO₂, and (D + N)eCO₂. In contrast, significantly higher shoot C concentrations ranked in AMF plants as (D + N)eCO₂ > DeCO₂ ≈ ACO₂ > NeCO₂. Meanwhile, only (D + N)eCO₂ increased root C concentrations in both non-AMF and AMF plants. However, neither shoot C nor root C concentrations were affected by the F. mosseae inoculation and by CO₂ × AMF interaction (Table 1).

Table 1. Carbon, nitrogen, phosphorus and potassium concentrations in shoots and roots of non-AMF and AMF-inoculated (AMF) wheat plants grown under ambient CO₂ (ACO₂) and under-elevated CO₂ concentrations at daytime (DeCO₂), night (NeCO₂), and both daytime and nighttime (D + N)eCO₂.

| Inoculation | CO₂ | Shoot | Root | Shoot | Root | Shoot | Root | Shoot | Root |
|-------------|-----|-------|------|-------|------|-------|------|-------|------|
| Non-AMF     |     |       |      |       |      |       |      |       |      |
| ACO₂        | 413 ± 10 a | 359 ± 10 b | 37.4 ± 1.8 a | 124 ± 1.1 ab | 8.39 ± 0.10 ab | 1.51 ± 0.07 ab | 31.49 ± 0.46 ab | 32.05 ± 0.06 ab |
| DeCO₂       | 445 ± 10 a | 401 ± 10 a | 26.0 ± 2.3 b | 9.5 ± 0.5 a | 8.80 ± 0.10 a | 0.96 ± 0.14 b | 29.66 ± 0.30 a | 6.72 ± 1.21 a |
| NeCO₂       | 430 ± 10 a | 393 ± 14 a | 27.6 ± 0.9 b | 12.6 ± 0.3 a | 5.15 ± 0.04 a | 1.17 ± 0.11 b | 23.60 ± 1.82 a | 8.56 ± 3.17 a |
| (D + N)eCO₂ | 448 ± 10 a | 431 ± 16 a | 28.5 ± 0.3 ab | 14.0 ± 1.0 a | 6.46 ± 1.34 ab | 1.38 ± 0.06 ab | 27.76 ± 0.76 a | 11.43 ± 1.94 a |

| AMF         |     |       |      |       |      |       |      |       |      |
| ACO₂        | 452 ± 7 b | 405 ± 15 b | 35.2 ± 0.7 a | 16.8 ± 0.9 a | 6.67 ± 0.55 a | 1.47 ± 0.13 a | 31.84 ± 0.61 a | 15.90 ± 0.63 a |
| DeCO₂       | 456 ± 8 b | 419 ± 7 b | 32.5 ± 1.2 a | 14.3 ± 0.9 a | 8.42 ± 0.63 a | 1.17 ± 0.16 a | 28.08 ± 1.06 a | 9.02 ± 1.15 a |
| NeCO₂       | 410 ± 11 c | 400 ± 16 a | 25.3 ± 0.6 c | 11.5 ± 1.5 b | 4.75 ± 0.47 c | 1.09 ± 0.15 c | 26.20 ± 1.64 a | 9.45 ± 0.98 a |
| (D + N)eCO₂ | 454 ± 12 a | 438 ± 10 a | 29.5 ± 1.4 b | 14.0 ± 0.8 b | 7.01 ± 0.49 b | 1.84 ± 0.19 a | 29.67 ± 0.82 a | 9.79 ± 0.72 a |

ANOVA

| CO₂        |     |       |      |       |      |       |      |       |      |
| AMF        | m   | n     | m    | *     | n    | m     | m    | n     | *    |
| eCO₂ × AMF | n   | m     | m    | *     | n    | m     | m    | n     | *    |

Data (means ± SE, n = 3) followed by different letters indicate significant differences between CO₂ treatments for the same AMF inoculation (a, b, c, d) and between AMF inoculations for the same CO₂ treatment (x, y) at p < 0.05. ANOVA: ns not significant; *, ** and *** indicate significant differences at p ≤ 0.05, p ≤ 0.01 and p ≤ 0.001, respectively.

Elevated CO₂ generally significantly decreased N in both the shoots and roots of AMF and non-AMF plants (Table 1), and such decreases were more pronounced under NeCO₂ in AMF plants, and DeCO₂ in non-AMF plants (Table 1). Reduction of shoot P concentrations were observed under both NeCO₂ and (D + N)eCO₂ in non-AMF plants and under NeCO₂ in AMF plants only. Both DeCO₂ and NeCO₂ decreased root P concentrations in both AMF and non-AMF plants. K concentrations in the shoot and root were generally significantly lower under eCO₂ than under ACO₂ in both AMF and non-AMF plants (Table 1). Meanwhile, F. mosseae inoculation significantly increased root N concentrations under both ACO₂ and DeCO₂, leaf N concentrations under DeCO₂ and root P concentrations under DeCO₂ and (D + N)eCO₂, and shoot K concentration under NeCO₂. A significant CO₂ × AMF interaction was observed in N concentrations in both shoots and roots, and the K concentration in roots, but not for P concentrations (Table 1).

3.3. Effects of AMF and CO₂ on Plant Biomass Production

Elevated CO₂ increased shoot, root and total plant biomass, regardless of whether the 12-week-old wheat seedlings were inoculated with F. mosseae or not (Figure 3A–C). Compared to the non-AMF plants, AMF colonization increased shoot and total plant biomass production, under ACO₂, DeCO₂ and (D + N)eCO₂, but not under NeCO₂ or for root biomass production (Figure 3B). Meanwhile, a significant CO₂ × AMF interaction was found for both the shoot and total plant biomass production (Figure 3A,C), but not for root biomass production (Figure 3B).
Figure 3. Effects of CO\(_2\) and mycorrhiza on (A) shoot, (B) root, and (C) total plant biomass production; (D) shoot C, (E) root C, and (F) total plant C accumulation; (G) shoot N, (H) root N, and (I) total plant N accumulation; (J) shoot P, (K) root P, (L) total plant P accumulation; and (M) shoot K, (N) root K, (O) total plant K accumulation in 12-week-old wheat grown under different daytime and/or nighttime CO\(_2\) concentrations inside environmentally controlled glass growth chambers. Values are the means ± standard error (SE), n = 3. Different letters above the bars indicate significant differences (p < 0.05), as revealed by Tukey’s test. Statistical comparisons (two-way ANOVA) between eCO\(_2\) or AMF treatments, as well as their eCO\(_2\) × AMF interaction are presented for each variable. Abbreviations: ACO\(_2\), ambient CO\(_2\) (410 ppm daytime + 460 ppm nighttime); DeCO\(_2\), elevated CO\(_2\) at daytime (550 ppm daytime + 460 ppm nighttime); NeCO\(_2\), elevated CO\(_2\) at nighttime (410 ppm daytime + 610 ppm nighttime); (D + N)eCO\(_2\), elevated CO\(_2\) at both daytime and nighttime (550 ppm daytime + 610 ppm nighttime). Daytime: 07:00 a.m.–19:00 p.m. and nighttime: 19:00 p.m.–07:00 a.m.
3.4. Effects of AMF and CO$_2$ on Plant C, N and P Accumulations

In general, eCO$_2$ significantly increased C accumulations in the shoot, root and total plant of the 12-week-old wheat seedlings (Figure 3D–F). The C accumulations in the shoot and total plant were significantly affected by *F. mosseae* colonization and CO$_2$ × AMF interaction (Figure 3D,F), but not in root C accumulations (Figure 3E). The accumulations of C in the shoot and total plant were significantly higher for the AMF plants under ACO$_2$, DeCO$_2$ and (D + N)eCO$_2$, but lower for the AMF plants under NeCO$_2$, compared to the respective non-AMF plants (Figure 3D–F).

Generally, eCO$_2$ significantly increased N accumulation in the shoot and total plant (Figure 3G,I), but not in the root (Figure 3H). Compared with non-AMF wheat plants, AMF plants had greater N accumulation in the shoot, root, and total plant under ACO$_2$, DeCO$_2$, and (D + N)eCO$_2$ (Figure 3G–I), but lower under NeCO$_2$ (Figure 3G,H). A significant CO$_2$ × AMF interaction on N accumulation was thus observed in the shoot, root, and total plant (Figure 3G–I).

Compared with ACO$_2$, eCO$_2$ significantly enhanced P accumulation in the shoot by 35–95%, and in the total plant by 30–79% in both non-AMF and AMF plants (Figure 3J,L), but had no effects on root P accumulation (Figure 3K). Neither the AMF symbiosis nor the CO$_2$ × AMF interaction showed a significant effect on the P accumulation in the shoot, root, and total plant (Figure 3J–L).

K accumulations in the shoot and total plant were significantly higher under all eCO$_2$ treatments in non-AMF plants, and under DeCO$_2$ and (D + N)eCO$_2$ in AMF plants, compared with ACO$_2$ (p < 0.05, Figure 3M,O). Significantly higher K accumulations in the shoot and total plant were in AMF than in non-AMF plants under ACO$_2$, DeCO$_2$, and (D + N)eCO$_2$. A significant CO$_2$ × AMF interaction on K accumulation was thus observed in the shoot and total plant (Figure 3M,O), but not in root K accumulations (Figure 3N).

3.5. Effects of AMF and CO$_2$ on Soil Nutrients

Soil AN, AP, and AK were significantly affected by CO$_2$, AMF, and CO$_2$ × AMF interaction, regardless of whether plants were colonized with AMF or not (Figure 4A–C). Compared with ACO$_2$, soil AN, AP, and AK were significantly decreased under all eCO$_2$ treatments in non-AMF soils, except AP under NeCO$_2$, and under DeCO$_2$ in AMF soils. Moreover, soil AN, AP, and AK were significantly increased by 5–36%, 7–49%, and 3–31% in AMF than in non-AMF soil under eCO$_2$ (Figure 4A–C).

3.6. Effects of AMF and CO$_2$ on Soil Enzymes Activity

In general, eCO$_2$ significantly increased the activity of soil invertase and urease (Figure 4D,E), but not the soil neutral phosphatase activity (Figure 4F) in both non-AMF and AMF plants. AMF colonization significantly increased the activity of invertase, urease, and neutral phosphatase (Figure 4D–F). Meanwhile, the CO$_2$ × AMF interaction only resulted in significant positive changes in the invertase activity (Figure 4D), but not in both the urease (Figure 4E) and neutral phosphatase activity (Figure 4F). In addition, the activity of invertase, urease, and neutral phosphatase was significantly increased by 15–56%, 19–50%, and 15–39% in AMF than in non-AMF soil under ACO$_2$, DeCO$_2$, and (D + N)eCO$_2$, but not under NeCO$_2$ (Figure 4D–F).

3.7. Correlations

Total plant biomass production was significantly negatively correlated to soil AN ($y = −0.10x + 12.08$, Figure 5A) and soil AK ($y = −0.06x + 17.42$, Figure 5C) in non-AMF plants, but not to AMF plants (Figure 5A,C). In contrast, no relationships between total plant biomass production and soil AP were observed in both AMF and non-AMF plants (Figure 5B).
Figure 4. Effects of mycorrhiza and CO$_2$ on (A) soil available nitrogen (AN), (B) soil available phosphorus (AP), (C) soil available potassium (AK), (D) invertase activity, (E) urease activity, and (F) neutral phosphatase activity in the soil of 12-week-old wheat grown under different daytime and/or nighttime CO$_2$ concentrations inside environmentally controlled glass growth chambers. Values are the means ± SE, $n = 3$. Different letters above the bars indicate significant differences ($p < 0.05$), as revealed by Tukey’s test. Statistical comparisons (two-way ANOVA) between AMF and CO$_2$ treatments, as well as their interaction (eCO$_2$ × AMF) are presented for each variable. Abbreviations are the same as in Figure 2.

Figure 5. Relationships between plant biomass production and (A) soil available nitrogen (AN), (B) soil available phosphorus (AP), (C) soil available potassium (AK), (D) soil invertase, (E) urease or (F) neutral phosphatase in the soil of 12-week-old wheat grown under different daytime and/or nighttime CO$_2$ concentrations inside environmentally controlled glass growth chambers. Data are means ± SE, $n = 12$. Regressions are shown for non-AMF (dotted lines) and for AMF (solid lines) treatments. Abbreviations are the same as in Figure 2.

In addition, total plant biomass production was significantly positively correlated to the invertase activity in both AMF ($y = 0.12x + 3.49$) and non-AMF plants ($y = 0.31x + 0.75$, Figure 5D), to the urease activity in AMF plants ($y = 1.30x + 2.44$), but not to non-AMF
plants (Figure 5E), and to the phosphatase activity in non-AMF plants \(y = 16.81x + 2.01\), but not to AMF plants (Figure 5F).

4. Discussion

4.1. Effects of AMF Symbiosis on Plant Biomass and C Accumulation Depend on eCO\(_2\) at Daytime or Nighttime

Studies on the understanding of mycorrhizal-CO\(_2\) responses are mostly focused on the differences between ACO\(_2\) and 550–1000 ppm eCO\(_2\) at daytime [17,19,55]; no information about mycorrhizal plants responses to eCO\(_2\) at nighttime has been reported. As a substrate for plant photosynthesis, eCO\(_2\) during daytime facilitates CO\(_2\) assimilation processes by increasing intercellular CO\(_2\) and leaf carboxylation efficiency of ribulose-1,5-bisphosphate carboxylase/oxygenase (RubisCO) while reducing photorespiration [6,56,57], leading to an accumulation of non-structural carbohydrates and a stimulation of biomass production (Figure 3A–F). Meanwhile, a part of the photosynthetically-fixed C is consumed by leaves, shoots, and roots through dark respiration [58]. As a product of plant respiration, an increased intercellular CO\(_2\) and reduced stomata conductance under nighttime eCO\(_2\) [59] would lead to a decrease of plant dark respiration [48]. As a result, more biomass production was found in wheat grown under 410/610 ppm NeCO\(_2\) than under 410/460 ppm ACO\(_2\) (Figure 3A–F), in line with earlier findings about Alfalfa [60], Phaseolus vulgaris [46], soybean [47], and Xanthium strumarium [44,47]. The decreased respiration was a physiological mechanism behind the increase in biomass by C conservation under NeCO\(_2\) [41,48,61].

Intriguingly, the most significant findings were that the pattern of plant response to eCO\(_2\) during daytime or nighttime was influenced by AMF inoculation. Plant growth depressions occur when increased nutrient benefits are outweighed by its C cost, whereas positive growth responses occur where benefits outweigh the cost [62]. The positive growth responses to AMF inoculation were observed under 550/460 ppm DeCO\(_2\) and 550/610 ppm (D + N)NeCO\(_2\), whereas a negative effect under 410/610 ppm NeCO\(_2\) on wheat growth and C accumulation was found in the present study (Figure 3A–F). With low daytime CO\(_2\) and high nighttime CO\(_2\), photosynthetic responses did not predominately control plant growth [43]. AMF colonization would drain more photo-assimilates from the host plant for extraradical hyphal growth [25] and increase mycorrhizal respiration under eCO\(_2\) than under ACO\(_2\) [37,63], leading to reduced pools of nonstructural carbohydrates in the host and growth depression under NeCO\(_2\) (Figure 3A–F). With high CO\(_2\) concentration at daytime, plant C utilization by AMF might be compensated by higher photosynthesis in host plants [64]. AMF would enhance more photo-assimilate production than that they could drain from the host plant due to an improved nutrient uptake, leading to a positive effect under DeCO\(_2\) and (D + N)eCO\(_2\) (Figure 3A–F), which is in line with that reported by Zhu et al. [65], who concluded that AMF (Rhizophagus irregularis)-colonized wheat achieved greater growth and higher C accumulation than non-AMF wheat at 700 ppm (D + N)eCO\(_2\) [65]. Thus, resource limitation is a key factor in the cost–benefit analysis of AMF effects on plant growth under NeCO\(_2\).

4.2. Nitrogen Demands are Increased under eCO\(_2\), but AMF Symbiosis Lessens N Limitation under eCO\(_2\) at Daytime, Not at Nighttime

Our results showed that eCO\(_2\) at daytime and/or nighttime caused a decrease in N concentrations of the shoot and root both in non-AMF and AMF wheat (Table 1), which was similar to earlier reports that tissue N concentrations were often decreased in wheat cultivars under 550–800 ppm eCO\(_2\) [25,32,65–67]. The following mechanisms could explain the decreased N in plants under eCO\(_2\) both at daytime and nighttime: (1) a “dilution effect” due to higher plant biomass production [9,68]; (2) reduced transpiration rates under eCO\(_2\) both at daytime and nighttime could decrease the transpiration-driven mass flow of nutrients, and hence, induced limitations in leaf nutrient transport led to decreased N uptake [66,69,70]; and (3) the reduction of the RubisCO protein that constitutes about half of the protein in leaves under eCO\(_2\) both at daytime and nighttime [48,71]. Beyond that, we speculated that the inhibition of leaf N assimilation under eCO\(_2\) was associated
with the reduction in respiration. The reduced respiration under high CO$_2$ has led to a reduced supply of energy-rich compounds, including ATP and NADH in the cytoplasm, and thereby decreases the amount of reductant available for NO$_3^-$ reduction [72,73].

The N concentrations of shoots and roots and the P concentration of roots were higher in AMF plants than in non-AMF counterparts grown under DeCO$_2$, whereas they were not affected by AM fungal colonization under NeCO$_2$ (Table 1). NeCO$_2$ hence resulted in a decrease in beneficial effects of AMF on plants. An enhanced C fixation under DeCO$_2$ would require more supply of N, P and K, leading to a strong decline in soil AN, AP, and AK associated with DeCO$_2$ (Figure 4A–C). Thus, _F. mosseae_ colonization promoted plant uptake of N, P, and K, and alleviated plant nutrient demands and soil N, P, and K limitations under eCO$_2$ at the daytime. In contrast, eCO$_2$ at nighttime decreased the dark respiration, resulting in reduced energy and nutrient demands [44]. In general, plants in nutrient-rich soils under NeCO$_2$ (Figure 4B,C) tend to be less frequently affected by AM fungi [74].

Although plant N concentrations were decreased due to CO$_2$ elevation, N accumulation in wheat tissues was enhanced because of an increased biomass in both non-AMF and AMF plants (Figure 3G–I). Similar to our findings, compared to non-AMF plants, leaf N% or plant total N accumulation under 700–1000 ppm eCO$_2$ was increased by 30–41% in AMF colonized wheat [65], alfalfa [36], and _Taraxacum officinale_ [31]. The significantly higher N in AMF plants might be due to a higher C accumulation from photosynthesis and hence a greater N demand under daytime eCO$_2$. In contrast, we found a negative AM effect on plant N uptake under nighttime eCO$_2$ (Figure 3G–I). One possible explanation could be that eCO$_2$ inhibited the assimilation of NO$_3^-$ [72], which is the dominant inorganic N form in dryland soils. The NO$_3^-$ assimilation of _Arabidopsis_ and wheat was slowed down under 720 ppm nighttime eCO$_2$ [72]. Thus, both the N form and atmospheric CO$_2$ concentration at nighttime are important factors in determining plant performance.

### 4.3. P and K Demands were Increased under eCO$_2$ at Daytime, Not at Nighttime

Shoot and root P or K concentrations were decreased under eCO$_2$, regardless of AMF status (Table 1). These results agreed with how tissue P and K concentrations in various legumes and non-legumes were often lower under 550–800 ppm eCO$_2$ than under ACO$_2$ because of an increase in dry matter and carbohydrate accumulation [4,25,75,76]. Such lower P or K concentrations in plant tissues could be alleviated by AMF colonization in some, but not in all cases. For example, leaf and root P concentrations in non-mycorrhizal _T. repens_ declined by 31% and 115% under 700 ppm eCO$_2$, whereas in _F. mosseae_ seedlings, the decline in P concentration was as low as 17% and 0.5%, respectively [77]. _R. intraradices_ stimulated the growth and P acquisition of sour orange, but not of sweet orange grown at high P (2 mM) supply under 700 ppm eCO$_2$ [37]. The growth of _Medicago truncatula_ and _Brachypodium distachyon_ under 900 ppm eCO$_2$ was increased by sufficient P supply, rather than by _R. irregularis_ colonization [9]. eCO$_2$ at 700 ppm resulted in a 20% or 22% decrease of K in grains of non-mycorrhizal or mycorrhizal durum wheat [33]. In general, plants have developed specific P acquisition strategies by root systems and/or mycorrhizal associations to take up limited P in the soil [78–80]. Plant P and K acquisition is obviously enhanced by extensive root development. The P and K concentration influences plant photosynthesis and growth rates, leading to multiple C–P and C–K interactions 550 ppm DeCO$_2$ (Figure 3J–O) while 610 ppm NeCO$_2$ did not increase shoot, root, or total plant P and K content (Figure 3J–O). The different response to DeCO$_2$ and NeCO$_2$ may be related to photosynthesis. As a general rule, AMF symbiosis delivered soil N, P, and K to plants in return for photosynthate, alleviating plant N, P, and K demands, resulting in increased growth responses to eCO$_2$ via positive feedback [3]. Thus, AMF symbiosis might alter the imbalance of sink and source under eCO$_2$ by regulating the demand for C and supply of N, P, and K from the soil to the host plant [35]. However, our results showed that AMF colonization had no significant effects on P accumulations in the shoot and root and K accumulation in the root (Table 1, Figure 3J–L,N). The mechanisms affecting P, K absorption
4.4. AMF Colonization Increased Soil N, P and K availability, Especially under eCO₂ at Daytime

Plant–soil interactive feedback to eCO₂ likely determines the increase or decrease in soil N, P, and K pools, as well as microbial community composition and activities. In general, C flows into the soil through the plant root and/or mycorrhiza directly or indirectly. Under eCO₂, more C was available for mycorrhizal growth and development, and AMF associations were thus stimulated by such extra C [81]. In turn, AMF accelerates soil organic matter decomposition under eCO₂ to mine for N and P [19]. The increase of invertase and urease activity under 500–610 ppm eCO₂ indicated higher decomposition of organic C-N-compounds and release of N and other nutrients, resulting in a general activation of microbes [82]. The relative increases in soil enzyme activities also might be attributed to a higher biomass production under eCO₂ (R² = 0.46–0.77, p < 0.001, Figure 5A,D–F), and hence a greater demand of N, P, and K [3]. In fact, the considerable decrease in available N, P, and K under DeCO₂ in the present study supports this view. A meta-analysis showed that eCO₂-induced nutrient limitation could increase soil enzyme activities [83]. However, a significantly higher level of soil AN, AP and AK was observed in AMF wheat seedlings than in non-AMF counterparts under eCO₂ (Figure 4A–C), suggesting that the presence of AM fungi could confer better soil fertility under eCO₂.

5. Conclusions

The present study under auto-simulated future daytime and/or nighttime eCO₂ provides evidence that eCO₂ can impact plant–soil feedback and/or plant–AMF symbiosis. Our results showed that the responses of crops to an averaged sole daytime/nighttime eCO₂ might not provide the expected effects of rising CO₂ concentration that they could have on crop growth. More soil N, P, and K nutrients were required under DeCO₂ to match increased C assimilation, leading to lower soil N, P, and K availability. Although F. mosseae colonization alleviated soil nutrient constraints in response to eCO₂, its role on plant growth depended on eCO₂ at daytime and/or nighttime, which could induce an imbalance in the source–sink relationship associated with reduced plant and soil N, P, and K. AMF symbiosis could improve plant C accumulation, N, P, and K uptake particularly under DeCO₂, while NeCO₂ decreased AMF’s beneficial effects on plants. Such information is essential for understanding the mechanisms influencing C, N, P, and K dynamics in future climate-change scenarios. As a result, integrating AMF’s benefits to plants under a factual field DeCO₂ and NeCO₂ will be critical when dealing with the long-term consequence of future CO₂ rising on global cropping systems.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/jof7060458/s1, Figure S1: Average daily (A, 417 ± 16/463 ± 27 ppm, means ± SD, n = 175, daytime (7:00 a.m.–19:00 p.m.)/nighttime (19:00 p.m.–7:00 a.m.)) and weekly (B, 421 ± 19/467 ± 18 ppm, means ± SD, n = 25, daytime/nighttime) variations of atmospheric CO₂ concentrations between October 2017 and March 2018 in the National Monitoring Base for Purple Soil Fertility and Fertilizer Efficiency (29°48′ N, 106°24′ E, 266.3 m above the sea level) on the campus of Southwest University, Chongqing, China. The atmosphere CO₂ concentrations were monitored by an auto-controlled facility (DSS-QZD, Qingdao Shengsen Institute of Science and Technology, Shandong, China). Figure S2: CO₂ auto-controlling facility (A) overview of environmentally controlled growth chambers, (B–C) CO₂ digital control screen, (D) the humidity, temperature, light and CO₂ sensors, (E–F) CO₂ gas cylinder, (G) the control system.

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