The Effects of Inorganic Nitrogen form and $\text{(CO}_2\text{)}$ Concentration on Wheat Yield and Nutrient Accumulation and Distribution

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The effects of inorganic nitrogen form and CO₂ concentration on wheat yield and nutrient accumulation and distribution

Eli Carlisle¹*, Samuel Myers², Victor Raboy³ and Arnold Bloom⁴

INTRODUCTION

Nitrogen (N) is the mineral element that most often limits plant growth and primary productivity in natural and agricultural systems. Plants usually acquire N from the soil in the forms of ammonium (NH₄⁺) and nitrate (NO₃⁻), and management of these forms is vital to agriculture. Wheat can utilize either form alone (Wang and Below, 1992), but mixed N nutrition (e.g., NH₄NO₃) typically produces the best grain yields and quality in hydroponically grown (Gentry et al., 1989; Heberer and Below, 1989; Wang and Below, 1995) and field-grown plants (Bock, 1987; Camberato and Bock, 1990).

Ammonium and nitrate affect crops differently when either is supplied as the sole N source (Bloom, 1997). Ammonium requires less energy to assimilate into organic compounds (Bloom, 1997), but can prove toxic if it accumulates to high concentrations within plant tissues (Cox and Reisenauer, 1973; Goyal and Huffaker, 1984). Nitrate is generally the predominant form available in aerated, temperate agricultural soils (Haynes, 1986; Bloom, 1997), and may accumulate within plant tissues to high concentrations without toxicity (Goyal and Huffaker, 1984). In wheat, the N form supplied has been found to influence many physiological parameters profoundly including biomass (Wang and Below, 1995, 1996, 1998; Bloom et al., 2002), leaf area (Bloom et al., 2002), tillering (Chen et al., 1998), seed mass (Wang and Below, 1995), protein content (Wang and Below, 1995), and mineral nutrient acquisition and distribution (Gashaw and Mugwira, 1981; Wang and Below, 1998), although such differences can vary among cultivars (Gashaw and Mugwira, 1981; Wang and Below, 1995).

The presence of NH₄⁺, as either a sole N source or in mixed N nutrition, increased organic N concentration in shoots, roots, and grain and decreased partitioning of dry matter to the roots in wheat (Wang and Below, 1995). Decreased cation uptake has been found in wheat under NH₄⁺ nutrition (e.g., Gashaw and Mugwira, 1981; Wang and Below, 1998), although results varied among cultivars (Gashaw and Mugwira, 1981). For example, NH₄⁺ nutrition decreased whole plant and shoot accumulations of K, Ca, Mg, Fe, Mn, and Zn (Wang and Below, 1998). Nutrient allocation to plant tissues also varied between N forms. NH₄⁺-fed plants distributed a smaller percentage of total P, K, Cu, and B to roots relative to NO₃⁻-fed plants (Wang and Below, 1995, 1998). Also, a greater percentage of reduced N was allocated to the shoots in NH₄⁺-fed plants (Wang and Below, 1995).

Elevated atmospheric concentrations of CO₂ alter growth and N dynamics of wheat and other C₃ plants. Under elevated CO₂, wheat has lower protein and N concentrations (e.g., Thompson and Woodward, 1994; Bloom et al., 2002; Wu et al., 2004), and lower macro- and micronutrients concentrations (Manderscheid et al., 1995; Fangmeier et al., 1997, 1999; Wu et al., 2004; Hög and Fangmeier, 2008). Grain phytate concentrations are also thought to increase or remain the same under elevated CO₂.
Cousins and Bloom, 2004) as well as in all other C$_3$ species grown to senescence. Here, we grew wheat hydroponically in an aerated nutrient solution that contained 1 mM CaSO$_4$, 1 mM K$_2$HPO$_4$, 1 mM KH$_2$PO$_4$, 2 mM MgSO$_4$, and 0.2 g L$^{-1}$ Fe–NaEDTA and micronutrients (20% of a modified Hoagland’s solution with either 0.2 mM KNO$_3$ or 0.1 mM (NH$_4$)$_2$HPO$_4$ as the N source, Epstein and Bloom, 2005). The nutrient solution was replaced weekly and an additional 0.2 mmoles of NO$_3$ - or NH$_4$ - N was added midweek until harvest. The solution volume was maintained by daily addition of deionized water. Solution pH varied between 6.8 and 7.0 for both of the N forms, and the NH$_4$ and the NO$_3$ solutions did not differ by more than 0.1 pH units.

The plants were grown in controlled environment chambers (Conviron, Winnipeg, Canada) set at 23/20°C day/night at 60–70% relative humidity with a photoperiod of 15 h. The photosynthetic flux density was 375 μmol m$^{-2}$ s$^{-1}$ at plant height. Plants were subjected to one of three CO$_2$ concentrations: “subambient” (310 ± 30 ppm), “ambient” (410 ± 30 ppm), and “elevated” (720 ± 50 ppm). Subambient CO$_2$ concentrations were maintained by passing air that entered the growth chamber through wet soda lime, a mixture of KOH, NaOH, and Ca(OH)$_2$ that was replaced as needed. The elevated CO$_2$ conditions were maintained in an environmental chamber equipped with non-dispersive infrared analyzers for CO$_2$ (Horiba model APRA-250E) and valves that added pure CO$_2$ to the incoming air stream to hold the chamber concentration at 720 ppm.

The wheat was grown until all aboveground parts turned completely yellow. Plant matter was sorted into grain, chaff, shoots, and roots and dried for 48 h at 55°C. Data on kernel number (KN), kernel mass, number of heads, kernels head$^{-1}$, and HI were collected prior to sample preparation for nutrient analysis. A portion of the grain was analyzed for phytate using a modification of the method as described by Haug and Lantzsch (1983). The remainder of the grain as well as the shoots and chaff was bulked into five repetitions per treatment and sent to the UC Davis Analytical Laboratory for nutrient analysis. The roots of plants for each CO$_2$ × N treatment became entangled within the same tub; therefore, we were unable to separate the roots of the individual plants for analysis. Root data are thus presented as means for each treatment with no standard errors or confidence intervals.

Data were analyzed using PROC MIXED (SAS 9.0 Cary, NC, USA). Nitrogen form and CO$_2$ factors were treated as fixed independent variables. We used the Tukey–Kramer Honestly Significant Difference test for mean separation. Probabilities less than 0.05 were considered significant. Because some of the transformed variables did not meet the assumption of homogeneity
of variances, but one-way ANOVAs met the ANOVA assumptions, we analyzed the results via one-way ANOVAs to gain some information on the interactions between CO₂ and N form.

**MODELING THE INFLUENCE OF N FORM ON Zn NUTRITION IN THE HUMAN DIET**

We used a database derived from the United Nation’s Food and Agriculture Organization (FAO)’s national food balance sheets (FBS) to estimate the average daily per capita dietary intake of zinc and phytate from 95 different food commodities in each of 176 countries. This database combines FAO data on per capita intake of food commodities with USDA data on the nutrient or phytate content of each of these commodities. More detailed discussion of the creation of this database for the International Zinc Collaborative Group may be found in Wuehler et al. (2005). Using this database, we produced two datasheets: one containing per capita daily dietary intake of zinc from each food commodity for each country and another containing per capita phytate intake from each food commodity for each country. To calculate total dietary zinc (TDZ) and total dietary phytate (TDP) per country, we summed across the rows of all food commodities for each respective country.

To determine the proportion of a population at risk for zinc deficiency from a hypothetical least developed country (LDC), we first calculated TDP and TDZ values for a set of 44 countries defined by the United Nations as being least developed. We took the mean TDP and TDZ values for these countries to represent a hypothetical “least developed country.” To calculate the bioavailable zinc portion (TAZ; usually a small fraction of TDZ) we used the Miller equation (Equation 1: Miller et al., 2007).

\[
TAZ = 0.5 \cdot \left( A_{\text{max}} + TDZ + K_R \cdot \left( 1 + \frac{TDP}{K_P} \right) \right) - \sqrt{\left( A_{\text{max}} + TDZ + K_R \cdot \left( 1 + \frac{TDP}{K_P} \right) \right)^2 - 4 \cdot A_{\text{max}} \cdot TDZ}
\]

**RESULTS**

We divide the results here into three categories: first, biomass and yield data for the shoots, grain, and roots; second, tissue concentrations and whole plant micro- and macronutrient contents; and third, nutrient distribution among the different tissues. Values of the statistical significance of the results were place into a table (Table 1) in order to improve the readability of the text.

**BIOMASS AND YIELD**

Plants supplied NH₄⁺ vs. NO₃⁻ nutrition reacted differently to CO₂ enrichment (Figure 1; Table 1). Plants supplied NH₄⁺ differed across CO₂ treatments for most of the yield and biomass measurements. The greatest values typically were found at ambient CO₂ concentrations. Shoot, chaff, grain yield, number of heads, and KN were greatest at ambient CO₂ levels. Individual kernel mass was greatest under both ambient and elevated CO₂ treatments. HI and kernels head⁻¹ showed no change across CO₂ treatments. In contrast, biomass and yield measures of NO₃⁻-supplied plants did not differ among the three CO₂ concentrations.

At subambient CO₂ differences between the NH₄⁺ and NO₃⁻ treatments occurred in shoot biomass and three of the yield components: kernel mass, head number, and kernels head⁻¹. Ammonium-supplied plants had a larger number of heads while NO₃⁻-supplied plants had greater shoot biomass, kernel mass, and kernels head⁻¹. At ambient CO₂, NH₄⁺-supplied plants had a greater number of heads and greater chaff biomass. Plants supplied NO₃⁻ had a larger number of kernels head⁻¹. At elevated CO₂, biomass and yield measures did not differ with N treatment.

**ROOT**

Roots had a smaller mean biomass when supplied NH₄⁺ than when supplied NO₃⁻ at all CO₂ concentrations (Figure 1). Both N treatments had the greatest biomass at ambient CO₂ and the smallest at subambient CO₂. The highest root:shoot ratios for both NH₄⁺ and NO₃⁻-supplied plants were observed at ambient and elevated CO₂. Ammonium-supplied plants always had lower root:shoot ratios and biomass than NO₃⁻-supplied plants at the same CO₂ concentration.

**NUTRIENTS**

**Total plant nutrients**

Total plant nutrients generally followed the same trend within N form, although NH₄⁺-supplied plants exhibited a greater diversity of responses to increasing CO₂ concentrations (Table 2). Total plant P, K, B, Ca, Mg, and Zn decreased with increasing CO₂ under NH₄⁺, while S and Mn were highest under ambient CO₂.
Table 1 | Results of a series of one-way ANOVAs run on the data.

<table>
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<td>NH₄⁺ NO₃⁻ NH₄⁺ NO₃⁻</td>
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<tr>
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<td></td>
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<td>P</td>
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<td>K</td>
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<td>Ca</td>
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<td>Mg</td>
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<td>Fe</td>
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<td>Cu</td>
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<td>NO₃⁻ – N</td>
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Among CO₂ conc. or between N forms

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<td>* NS</td>
<td>NO₃⁻</td>
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<td>NS NS</td>
<td>NO₃⁻</td>
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</table>

Differences among CO₂ concentrations within an N form and between N forms at each CO₂ concentration for shoot and grain nutrient concentrations. Differences among CO₂ concentrations within an N form or between N forms at each CO₂ concentration for biomass and yield components. The symbols indicate statistical significance (*0.05, **0.01, ***0.001, NS, not significant).

Ammonium-supplied plants had the greatest amounts of Fe and total N at subambient CO₂. Nitrate-supplied plants accumulated the greatest amounts of total N, P, K, S, B, Ca, Zn, Mn, and Mg at ambient CO₂. Only three nutrients – K, S, and Fe – had the lowest contents at elevated CO₂.

Shoot

Under NH₄⁺ supply, plants varied with CO₂ concentration for total N, P, S, Ca, Cu, B, Mn, Zn, and NO₃⁻ – N (Table 1; Figure 2). Calcium and Cu were highest under subambient CO₂. Total N and S were greatest at subambient and elevated CO₂. Nitrate-N was greatest at ambient CO₂. Phosphorus was highest at elevated CO₂ concentrations. Boron, Zn, and Mn increased with CO₂ concentration.

Plants supplied NO₃⁻ showed significant variation across CO₂ treatments for K, Ca, Mg, B, Fe, Cu, Zn, and NO₃⁻ – N (Table 1; Figure 2). Calcium and Cu had the greatest concentrations at subambient CO₂. The highest concentrations of B, Fe, and Zn occurred at subambient and elevated CO₂. Potassium concentrations were highest at elevated CO₂. Nitrate-N increased with CO₂. Magnesium showed the opposite trend, decreasing with CO₂ concentration.

Differences between N forms were also evident. At subambient CO₂, NH₄⁺-supplied plants had increased concentrations of P, S, and Zn, while NO₃⁻-supplied plants had greater concentrations of B, Mg, Mn, and NO₃⁻ – N (Table 1; Figure 2). Concentrations of K, Zn, and Cu were higher in plants supplied NH₄⁺ at ambient CO₂, while Mg, Mn, and NO₃⁻ – N were greater in plants supplied NO₃⁻. At elevated CO₂, concentrations of N, P, S, and Zn were higher in plants supplied NH₄⁺, while concentrations of B, Mg, Mn, and NO₃⁻ – N were greater in plants supplied NO₃⁻.

Grain

Grain nutrient concentrations. Plants supplied NH₄⁺ showed significant variation across the CO₂ treatments in the concentrations of total N, P, K, Ca, B, Cu, Mn, and NO₃⁻ – N (Table 1; Figure 3). The greatest concentrations of total N, P, K, Ca, and Cu were found at subambient CO₂. Iron concentrations were high at both subambient and ambient CO₂. Boron was equally high at subambient and elevated CO₂. Manganese was greatest at elevated CO₂. Nitrate-N decreased with increasing CO₂.

Significant differences among the NO₃⁻-supplied plants across CO₂ treatments were only observed in S and B. The greatest concentrations of B were found at subambient CO₂. Sulfur was highest at ambient CO₂.

Nitrogen form significantly affected grain nutrient concentrations (Table 1; Figure 3). At subambient CO₂, NH₄⁺-supplied plants had higher concentrations of total N, K, S, Ca, Zn, and Cu than NO₃⁻ plants. At ambient CO₂, Ca, Zn, and Cu were greatest under NH₄⁺. Ammonium-supplied plants also had the highest concentrations of K, S, Ca, Zn, and Cu at elevated CO₂. At no CO₂ concentration did plants supplied NH₄⁺ have significantly lower concentrations of any micro- or macronutrient than those supplied NO₃⁻.

Phytate and bioavailable Zn. Phytate was relatively insensitive to CO₂ concentration. Phytate concentrations were highest...
at subambient CO\textsubscript{2} for NH\textsubscript{4}\textsuperscript{+}-supplied plants (Figure 4). Sub-ambient CO\textsubscript{2} also produced the lowest phytate concentrations in NO\textsubscript{3}\textsuperscript{-}-supplied plants. NH\textsubscript{4}\textsuperscript{+}-supplied plants had greater phytate concentrations than NO\textsubscript{3}\textsuperscript{-}-supplied plants at subambient CO\textsubscript{2}, but not at the other CO\textsubscript{2} concentrations. Grain from plants grown under NH\textsubscript{4}\textsuperscript{+} nutrition had roughly 7, 18, and 8% higher bioavailable Zn than NO\textsubscript{3}\textsuperscript{-}-supplied plants at subambient, ambient, and elevated CO\textsubscript{2}, respectively (Figure 4).

Based on this phytate and bioavailable Zn data, we modeled how a human population from a LDC would be affected by
changes in atmospheric CO$_2$ concentrations (Table 3). The calculations were based on differences among CO$_2$ concentrations; therefore, modeled TDZ, TDP, and TAZ values cannot be compared between NH$_4^+$ and NO$_3^-$-supplied grain. Grain from plants supplied the different N forms behaved differently as CO$_2$ concentration increased. We found that under NH$_4^+$ supply, TAZ would increase 3.6% with the rise in CO$_2$ from subambient to ambient, and decrease 1.6% with the rise from ambient to elevated CO$_2$ (Figure 4). Humans provided NO$_3^-$-supplied wheat would experience a decrease in TAZ of 3.5% going from subambient to ambient, and an increase 5.6% from ambient to elevated CO$_2$ (Figure 4).

**Roots**

Ammonium-supplied plants generally showed a trend toward decreasing nutrient concentrations with increasing CO$_2$ concentration while NO$_3^-$-supplied plants varied widely across CO$_2$ treatments (Figure 5). The decrease in nutrient concentrations under NH$_4^+$ supply corresponded to an increase in root mass. Nitrate-supplied plants tended to have their highest nutrient concentrations in the ambient and elevated CO$_2$ treatments. Ammonium-supplied plants had higher concentrations of Zn and Mn across all of the CO$_2$ treatments, as well as higher total N and Fe at subambient CO$_2$. Nitrate-supplied plants typically had higher concentrations of the other nutrients at all CO$_2$ concentrations.

**Distribution of nutrients**

The distribution of nutrients and micronutrients among plant parts followed similar patterns in both the NH$_4^+$ and NO$_3^-$-supplied plants, although the NH$_4^+$-supplied plant distributions were slightly more variable (Table 4). Allocations to root and grain usually were greatest at ambient CO$_2$, and those to shoot at either subambient or elevated CO$_2$. Grain typically contained the largest proportion of total N, P, Zn, and Cu, although the organ with the largest percentage of Cu varied with CO$_2$ treatment among NO$_3^-$-supplied plants. Plants at subambient and elevated CO$_2$ allocated more Cu to the grain, while those at ambient CO$_2$ allocated more to the roots. In general shoots received the majority of K, S, B, Ca, and Mg for all N and CO$_2$ treatments. Ammonium-supplied plants allocated slightly more Mn to the roots at subambient CO$_2$, but allocated increasing amounts to the shoots at the expense of the roots as CO$_2$ concentration increased. In contrast, NO$_3^-$-supplied plants allocated most of the Mn to the shoots. Ammonium-supplied plants typically allocated more resources to the chaff while NO$_3^-$-supplied plants allocated a greater percentage of elements to the roots.

**DISCUSSION**

No other study to our knowledge has examined the influence of N form (NH$_4^+$ vs. NO$_3^-$) on plant nutrient relations at three different atmospheric CO$_2$ concentrations. Overall, N form affected growth, total plant nutrient contents, and nutrient distribution in senescing wheat shoots, grain, and roots. The influence of NH$_4^+$ and NO$_3^-$ on growth and nutrient status were so distinct that they should be treated as separate nutrients and not bundled into a general category of N nutrition. Wheat size and nutrition at senescence responded to CO$_2$ concentration in a non-linear manner. As was previously shown (Bloom et al., 2012), we found that plants supplied with NH$_4^+$ were more responsive to CO$_2$ concentration than those supplied with NO$_3^-$.

Although not explicitly addressed here because of the heterogeneity of variances, interactions between CO$_2$ and N treatments likely existed for a number of the biomass and nutrient measures. Most nutrient concentrations were generally higher in NH$_4^+$-supplied plants, with the exceptions of NO$_3^-$ – N, Mg, B, and Mn, which were generally higher in NO$_3^-$-supplied plants. Phytate, which hinders human absorption of Zn and Fe (Raboy, 2009), showed little variation at ambient and elevated CO$_2$ between NH$_4^+$ and NO$_3^-$-supplied plants, which, in conjunction with the observed greater bioavailable of Zn in NH$_4^+$-supplied plants, may have consequences for human nutrition. Distribution of nutrients to the shoots, roots, and grain in response to CO$_2$ concentration and N form was also non-linear and varied by nutrient.

**BIOMASS AND YIELD**

The data support our hypothesis that NO$_3^-$-supplied plants would show a more limited biomass and yield enhancement with CO$_2$ enrichment than NH$_4^+$-supplied plants. Nevertheless, mean biomass and yield decreased from ambient to elevated CO$_2$ in both NO$_3^-$- and NH$_4^+$-supplied plants in contrast to biomass increases in prior work on wheat seedlings (Bloom et al., 2002). NO$_3^-$-supplied plants allocated more biomass to roots and had larger root:shoot ratios than NH$_4^+$-supplied plants regardless of CO$_2$ concentrations as has been reported previously (Wang and Below,
FIGURE 2 | The effect of N form and CO₂ concentration on shoot nutrient concentrations of wheat grown hydroponically to senescence. Closed (NH₄⁺) and open (NO₃⁻) symbols represent back-transformed means and 95% confidence intervals (n = 5). Macro- and micronutrients are listed in the upper left of each frame. Differences are significant within N form if letters are different. Differences between N forms at each CO₂ concentration are generally significantly different if error bars do not overlap (see Table 1 for statistical significance).
FIGURE 3 | The effect of N form and CO$_2$ concentration on grain nutrient concentrations of wheat grown hydroponically to senescence. Closed ($\text{NH}_4^+$) and open (NO$_3^-$) symbols represent back-transformed means and 95% confidence intervals ($n = 5$). Macro- and micronutrients are listed in the upper left of each frame. Differences are significant within N form if letters are different. Differences between N forms at each CO$_2$ concentration are generally significantly different if error bars do not overlap (see Table 1 for statistical significance).
1995; Bloom et al., 2002), but increased root mass at elevated CO$_2$ concentration for NO$_3^-$-supplied plants reported previously (Bloom et al., 2002) were not observed here. The shoot biomass data suggest that growth differences measured early in the lifespan of wheat supplied with NH$_4^+$ or NO$_3^-$ or NH$_4^+$/NO$_3^-$ (i.e., greater shoot biomass in plants supplied NH$_4^+$ relative to those supplied NO$_3^-$ at elevated CO$_2$ concentrations; Bloom et al., 2002) do not necessarily carry through to senescence. This may be due in part to a shift in NO$_3^-$ assimilation to the root (Kruse et al., 2003), allowing NO$_3^-$-supplied plants to compensate for the decrease in shoot NO$_3^-$ assimilation that occurs at elevated atmospheric CO$_2$ concentrations (Bloom et al., 2002, 2010, 2012).

The decrease in yield and biomass measures at elevated CO$_2$ concentrations does not agree with field observations where wheat yields as well as overall biomass increased with elevated CO$_2$ (Högy and Fangmeier, 2008; Taub et al., 2008). Similarly, our results that the greatest values for other yield measures (e.g., heads, kernel mass, KN) occurred at ambient CO$_2$ concentrations vary from the literature. High CO$_2$ has been found to increase flowering tillers (Havelka et al., 1984; Fangmeier et al., 1996), KN (McKee et al., 1997), and kernel mass (i.e., thousand grain weight; McKee et al., 1997). Conflicting results, however, have also been reported (e.g., Havelka et al., 1984). Many of the field and open top chamber studies were grown under natural light and thus received substantially greater photosynthetic flux density than our chamber-grown plants. These higher light conditions would be more favorable to biomass accumulation. Also, these studies typically applied high amounts of mixed N fertilizer (e.g., NH$_4$NO$_3$), and yields and biomass have been found to be greater under mixed N nutrition than under either NH$_4^+$ or NO$_3^-$ alone (Cox and Reisenauer, 1973; Gentry et al., 1989; Heberer and Below, 1989; Wang and Below, 1995). Finally, the wheat cultivar we used (T. aestivum cv. Very 10) is a short-statured variety that has rarely been used in other studies and may have accounted for some of the differences between our study and other published data.

Our results that NH$_4^+$-supplied plants had greater yield and yield components than NO$_3^-$-supplied plants at ambient CO$_2$ have been observed previously (Wang and Below, 1996; Chen et al., 1998). Wang and Below (1995) observed greater numbers of kernels head$^{-1}$ and KN in plants supplied NO$_3^-$ that was not observed here. Their study, however, supplied NH$_4^+$ at relatively high levels (~8.9 vs. 0.2 mM NH$_4^+$ – N in our study). Several studies (Bennett and Adams, 1970; Cox and Reisenauer, 1973) have found that incipient NH$_4^+$ toxicity can start appearing at N levels as low as 0.08–0.2 mM NH$_4^+$, although the onset of NH$_4^+$ toxicity depends on light level (Magalhaes and Wilcox, 1984; Britto and Kronzucker, 2002) and solution pH (Findenegg, 1987). The poorer performance of the NH$_4^+$ treatment in Wang and Below (1995), therefore, might derive from NH$_4^+$ toxicity. We have previously determined that the 0.2 mM NH$_4^+$-supplied to our plants to be sufficiently high for normal growth, but low enough to avoid toxicity problems under our experimental conditions (Bloom et al., 2002).

**PLANT NUTRIENTS**

Our second hypothesis, that nutrient concentrations are differentially affected by the inorganic N form supplied to the plants and CO$_2$ enrichment, was supported by our data. CO$_2$ concentration...
FIGURE 5 | The effect of N form and CO₂ concentration on root tissue nutrient concentrations of wheat grown hydroponically to senescence. Closed (NH₄⁺) and open (NO₃⁻) symbols represent the bulked treatment mean (n = 10). Macro- and micronutrients are listed in the upper left of each frame. The lack of error bars reflects that the root mass for each treatment was bulked and analyzed as a unit.
and N form interactions may alter tissue demands for nutrients. For many nutrients, ratios between different elements are typically maintained within a narrow range (Garten, 1976; Bloom et al., 1985; Loladze, 2002). CO₂ concentration and N form may disturb the balance between different nutrients, leading to a cascade of changes in demand, accumulation, and allocation among the different plant tissues (e.g., Loladze, 2002; Högy and Fangmeier, 2008; Natali et al., 2009). Nitrate-supplied plants accumulated the greatest amounts of nutrients at ambient CO₂ (Table 2). Some portion of the greater response of NH₄⁺-supplied plants to CO₂ derived from a dilution effect from the greater biomass at ambient CO₂ concentrations (Figures 2 and 3). Total amounts of nutrients tended to decline with CO₂ enrichment for NH₄⁺-supplied plants, which had the greatest amounts of macro/micronutrients at subambient CO₂ (Table 2). These results have not been observed in other published studies (e.g., Fangmeier et al., 1997; Wu et al., 2004). Growth chamber studies, however, tend to have more exaggerated differences among treatments than field and greenhouse experiments (Högy and Fangmeier, 2008), and N source cannot be well-controlled in field and greenhouse experiments.

The observed increase in NO₃⁻–N concentration with CO₂ concentration in NO₃⁻-supplied plants has been reported previously (Bloom et al., 2002), and adds further support to the hypothesis that elevated CO₂ concentrations and the resulting decrease in photorespiration inhibit shoot NO₃⁻ photoassimilation. Nevertheless, tissue NO₃⁻–N concentrations observed here were substantially lower than those in the earlier study (Bloom et al., 2002). Again, this may derive from difference in life stages in the two studies. Most of the N available to the plant for grain filling comes from N translocation rather than uptake from the substrate (Simpson et al., 1983). Probably, the plants continued to assimilate plant NO₃⁻ using a non-photorespiratory dependent process such as root assimilation after root N uptake slowed or stopped. Loss of NO₃⁻ through root efflux to the nutrient solution also may have contributed to the lower concentration of NO₃⁻–N.

The partitioning and accumulation of all mineral elements was affected in some manner by the CO₂ treatment and N form supplied to the plants. Observations that cation concentrations decrease under NH₄⁺ supply (e.g., Cox and Reisenauer, 1973; Gashaw and Mugwira, 1981; Wang and Below, 1998) relative to NO₃⁻ supply were not apparent in this study. Again, this could be partly due to the relatively low concentration of NH₄⁺ supplied in our study, the age of the plants at harvest, and differences among wheat cultivars.

Allocation of nutrients within the plant followed similar trends for both N forms, with the exceptions of Mn and Cu (Table 2). Interestingly, in NO₃⁻-supplied plants, shoot Mn concentrations increased slightly with CO₂, and these plants allocated far more Mn to the shoots than NH₄⁺-supplied plants at all CO₂ concentrations. Manganese (Mn³⁺) has been found to activate Rubisco in place of Mg²⁺ and the Rubisco-Mn complex has been observed to decrease Rubisco carboxylase activity while minimally affecting or even enhancing oxygenase activity (Jordan and Ogren, 1983). The slight increase in shoot Mn with CO₂ corresponded to a large 23% decrease in Mg concentration. Manganese, which can act as a cofactor for glutamine synthetase (Smirnoff and Stewart, 1987), was also the only nutrient that NH₄⁺-supplied plants allocated a
greater percentage to the roots at the expense of the shoots. NO$_3^-$-supplied plants typically allocated a higher percentage of most nutrients to the roots, as has been reported previously (Wang and Below, 1995, 1998).

Phytate, which forms complexes with divalent cations, has been found to hinder human Zn and Fe absorption during digestion and thus has been labeled an “anti-nutrient.” It may serve a number of valuable functions, however, including roles as an anti-oxidant and anti-cancer agent (Raboy, 2009). Phytate is also the major repository of grain P, and variation in P supply to the developing seed is the major determinant of net seed phytate accumulation (Raboy, 1997, 2009; Cakmak et al., 2010). To our knowledge, no published studies have explicitly looked at how phytate is affected by CO$_2$ concentration. Elevated CO$_2$ has been found to have a much larger negative impact on Zn and Fe concentrations than on P in wheat (Loladze, 2002; Cakmak et al., 2010). Several studies (e.g., Fangmeier et al., 1999; Hogy and Fangmeier, 2008) have observed that P increases slightly with CO$_2$ concentration, and because the majority of P is tied up in phytate, this may cause increases in grain phytate concentrations as atmospheric CO$_2$ rises. As a result, bioavailable Zn and Fe–Zn and Fe not bound to phytate – is expected to decrease even further (Loladze, 2002).

Nonetheless, we did not observe such trends in macro- and micronutrient concentrations in this study. The mechanism behind these contrasting results is not clear, although the environmental conditions and nutrient solution in which the plants were grown likely had some role. The modeled data demonstrated only a small negative impact of CO$_2$ concentration on bioavailable Zn concentrations (Table 4), which was unexpected. Indeed, the grain from NO$_3^-$-supplied plants actually showed a slight increase in bioavailable Zn between ambient and elevated CO$_2$. These results combined with the differences in grain bioavailable Zn between NH$_4^+$ and NO$_3^-$-supplied plants demonstrates that N form may differentially affect the nutritional status of this important nutrient, especially in less developed countries that might be more dependent on phytate-rich grains for their Zn nutrition (Table 3). The milling process removes some, if not most, of the phytate and grain mineral content with the bran fraction of the grain (Guttieri et al., 2006). Regardless, with over 50% of the human population suffering from Zn deficiencies, even small increases in bioavailable Zn would be beneficial (Loladze, 2002). This modeling exercise, however, is not a prediction of how increasing CO$_2$ will affect wheat nutrition so much as illustrates that N source may mediate, to some extent, the effects of CO$_2$ on phytate and bioavailable Zn, and that N source will become an even more important agricultural consideration in the future.

In summary, both CO$_2$ concentration and N form strongly affect biomass and yield in hydroponically grown wheat, as well as nutrient concentrations in above- and belowground tissues. Interactions among plant nutrient concentrations, CO$_2$ concentrations, and N form are complex and non-linear. The impact of N form and CO$_2$ concentration on the mechanisms affecting nutrient accumulation and distribution requires further research and extension to more realistic and agriculturally relevant growing conditions found in greenhouse and field studies. Of course, in greenhouse and field studies, control of N source is limited and control of atmospheric CO$_2$ concentration is expensive. The effects of CO$_2$ and N form on agriculture and human nutrition observed here are interesting and suggest a new area of research on mitigating the effects of climate change on agriculture. The supply of fertilizers (e.g., urea, NH$_4$NO$_3$, anhydrous NH$_3$, organic amendments) or addition of nitrification inhibitors that increase the amount of available NH$_4^+$ may have beneficial effects for human nutrition, particularly in regards to micronutrient deficiencies such as Zn and Fe that currently affect billions of people worldwide. In the face of the potentially negative consequences of climate change on agriculture, all avenues of mitigation must be examined, and even small improvements may prove worthwhile.

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