Quantifying the Acidic and Basic Effects of Vegetable and Herb Species in Peat-based Substrate and Hydroponics

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Abstract. Objectives were 1) to quantify acidic and basic effects on the root zone pH for eight vegetable and herb species grown in peat-based substrate and hydroponic nutrient solution and 2) to determine the applied NH4+:NO3– ratio expected to have a neutral pH reaction for each species during its vegetative growth phase. In one experiment, plants were grown for 33 days in substrate (70% perlite:30% vermiculite by volume), and were fertilized with a nutrient solution containing 7.14 milli-equivalents (mEq)-L-1 N and NH4+:NO3– ratios ranging from 0:100 to 40:60. During the second experiment, the same species were grown in hydroponic nutrient solutions at 7.14 mEq-L-1 N with NH4+:NO3– ratios ranging from 0:100 to 30:70, and data were collected over a 6-day period. In substrate, species increased root zone pH when supplied 0:100 solution, except for cucumber, which did not change substrate pH. Increasing the NH4+:NO3– ratio to 40:60 increased acidity and decreased pH across species. Similar trends were observed in hydroponics, in which the most basic response occurred across species with 0:100, and the most acidic response occurred with 30:70. Arugula was the only species that increased root zone pH with all three NH4+:NO3– ratios in substrate and hydroponics. In substrate and hydroponics, mEq of acidity (negative) or basicity (positive) produced per gram dry weight gain per plant (mEq-g-1) correlated positively with mEq-g-1 net cation minus anion uptake, respectively, in which greater cation uptake resulted in acidity and greater anion uptake resulted in basicity. In hydroponics, the greatest net anion uptake occurred with 0:100, and increasing the NH4+:NO3– ratio increased total cation uptake across species. Cucumber had the most acidic effect and required less than 10% of N as NH4– for a neutral pH over time, arugula was the most basic and required more than 20% NH4–, and the remaining species had neutral percent NH4– between 10% and 20% of N. Increasing the NH4+:NO3– ratio decreased Ca2+ uptake across all species in hydroponics, which could potentially impact tip burn and postharvest quality negatively. Controlling root zone pH in substrate and hydroponic culture requires regular pH monitoring in combination with NH4+:NO3– adjustments and other pH management strategies, such as injecting mineral acid to neutralize irrigation water alkalinity or adjusting the limestone incorporation rate for substrate.

Nutrient availability for plant uptake is affected by root zone pH, especially in the case of metal micronutrients such as Fe2+ and Mn2+, which decrease in solubility as pH increases (Lindsay, 1979; Peterson, 1981). Several factors interact and affect root zone pH during soilless and hydroponic production, including N forms in the applied fertilizer, water alkalinity and pH buffering, and plant species (Bar-Yosef, 2008; Bugbee, 2004; Sonneveld and Voogt, 2009). Controlling root zone pH and preventing drift is, therefore, an important aspect of nutrient management. Plants affect root zone pH primarily through differential uptake of cation and anion nutrients (Haynes, 1990; Lea-Cox et al., 1996; Marschner, 2012; Rengel, 2003). Plants maintain a neutral charge across root membranes, compensating for a greater net uptake of either cations or anions by roots through endogenous efflux of H+ and OH–/HCO3– ions, respectively, into the root zone solution. Relative cation vs. anion uptake can be expressed as both a net difference of mEq, or as a ratio of cation-to-anion uptake (Kirkby and Knight, 1977; Rengel, 2003). Ratio values greater than one, less than one, and equal to one indicate greater cation uptake, greater anion uptake, and equal uptake, respectively. Net uptake of cations and anions has been correlated with the amount of acid and base produced in the root zone for crop species grown in field soil, soilless substrate, and hydroponic solutions (Dickson et al., 2016; Kirkby and Knight, 1977; Lea-Cox et al., 1996; Marschner, 2012; Rengel, 2003).

Nitrogen represents ≈70% to 80% of total nutrient uptake and can be taken up in cationic (NH4–) or anionic (NO3–) form, and therefore has a major impact on root zone pH (Lea-Cox et al., 1999; Marschner, 2012). Supplying NH4– results in root zone acidity from greater net cation uptake and efflux of H+ ions following uptake of NH4– by roots (Haynes, 1990; Kirkby and Knight, 1977; Lea-Cox et al., 1999; van Beusichem et al., 1988). A supply of NO3– results in root zone basicity from OH–/HCO3– efflux following uptake by roots (Haynes, 1990; Kirkby and Knight, 1977; Lea-Cox et al., 1999; van Beusichem et al., 1988). Uptake of NH4– is favored energetically over NO3– when both forms are supplied (Engels and Marschner, 1995). Conversion of NH4– to NO3– by microbial nitrification also produces H+ ions and root zone acidity, affecting root zone pH in both soilless substrate (Lang and Elliot, 1990) and recirculating hydroponic systems (Bugbee, 2004; Scharzw et al., 1999).

The effect of nutrient solution NH4+:NO3– ratio on root zone pH has been studied in greenhouse vegetable species grown in hydroponic culture (Bar-Yosef, 2008; Conesa et al., 2009; Ikeda and Osawa, 1983; Imas et al., 1997; Suvvas et al., 2006; Sonneveld and Voogt, 2009). Edible crop species are also grown in peat-based soilless substrate for container crop and food production (Hamrick, 2003; Resh, 2001). Sonneveld and Voogt (2009) and Bar-Yosef (2008) report that adjustment of the ratio of N forms and the total N concentration are strategies for managing root zone pH in hydroponic nutrient solution and soilless substrates. However, there is limited information comparing the pH effects of multiple vegetable and herb species, which would be useful in developing fertilizer and pH management strategies.

The objectives of this study were 1) to quantify the effects on root zone pH for eight vegetable and herb species grown in peat-based substrate and hydroponic nutrient solution, and 2) to determine specific NH4+:NO3– ratios for each species estimated to have a neutral pH effect in both substrate and hydroponic conditions during the vegetative growth stage. Species pH effect was...
could require a greater NH₄⁺/NO₃⁻ ratio to stabilize root zone pH.

**Materials and Methods**

*Substrate experiment.* A factorial experiment using a randomized complete block design evaluated eight species fertilized with three nutrient solution NH₄⁺/NO₃⁻ N ratios (0:100, 20:40, and 40:60) from a 0.5x modified Hoagland’s solution supplied at 7.14 mEq L⁻¹ N for effects on root zone acidity–basicity. Seedlings plugs of eight edible species were transplanted from 144-cell trays (Knox Nursery, Winter Garden, FL) into four-cell plastic bedding plant containers (88 mL/plug or 352 mL/container) at one plant per cell (four plants per container). There were six replicates per treatment, where each replicate was one four-cell container. Species consisted of arugula (*Eruca sativa* Mill.), basil (*Ocimum basilicum* L., ‘Genovese’), bell pepper (*Capsicum annuum* L., ‘Yankee’), cucumber (*Cucumis sativus* L., ‘Marketmore 76’), eggplant (*Solanum melongena* L., ‘Galine F1’), lettuce (*Lactuca sativa* L., ‘Green Star Green Grand Rapids’), spinach (*Spinacia oleracea* L., ‘Tyee F1’), and tomato (*Solanum lycopersicum* L., ‘Big Beef’)

The substrate was 70% peat/30% perlite (v/v) using Canadian sphagnum peat (Sun Gro Horticulture, Bellevue, WA) with long fibers and little dust [von Post scale, 1–2 (Puustjarvi and Robertson, 1975)]. Hydrated dolomitic limestone [Graymont Western Lime, Inc., West Bend, WI; 97% Ca(OH)₂–MgO, of which 92% passed through a 45-μm mesh and had an acid neutralizing value of 140% calcium carbonate equivalents (CCE)] was incorporated at 1.67 kg m⁻³ substrate to adjust pH to 6.0. Plants were grown on benches in a controlled-environment growth chamber located at the University of Florida in Gainesville, FL, for 33 d. Lighting was supplied by cool-white fluorescent bulbs (32 W, 6500K) at 175 μmol m⁻² s⁻¹ with an 18-h photoperiod that provided a daily light integral of 11.3 mol m⁻² d⁻¹ photosynthetically active radiation. Daily air and substrate temperatures during the experiment were 22.7 ± 1.5 °C and 22.3 ± 1.6 °C (mean ± se), respectively.

Modified nutrient solutions (0.5x Hoagland’s nutrient solution) were mixed with reagent-grade salts in zero alkalinity deionized water. Macronutrients were supplied at (mEq L⁻¹) 7.14 N, 0.52 P, 2.99 K, 5.00 Ca, and 1.00 Mg. Sulfate with NH₄⁺/NO₃⁻ ratio because NH₄⁺-N was supplied from ammonium sulfate, and S concentration was 2.38, 3.80, and 7.06 mEq L⁻¹ for the 0:100, 20:80, and 40:60 solutions, respectively. Macronutrients were derived from (NH₄)₂SO₄, Ca(NO₃)₂·4H₂O, KH₂PO₄, MgSO₄·7H₂O, CaCl₂·2H₂O, KNO₃, and K₂SO₄. Micronutrient concentrations were constant at (μEq L⁻¹) 72 Fe, 36 Mn, 6 Cu, 15 Zn, and 1 Mo derived from FeEDDHA (6.0% Fe; Akzo-Nobel, Holland); CuSO₄·5H₂O, MnSO₄·H₂O, (NH₄)₂MoO₄·2H₂O, and ZnSO₄·7H₂O. The B form is primarily as an uncharged molecule at pH less than 7 (Marschner, 2012) and was supplied at 1.0 mg L⁻¹ from H₂BO₃.

Replicate containers were irrigated at transplant to container capacity with 150 mL 0:100, 20:80, or 40:60 solution at 14.28 mEq L⁻¹ N, and each subsequent irrigation consisted of 100 mL replicate at 7.14 mEq L⁻¹ N. A plastic liner with a flat bottom was placed under each replicate to allow for leachate collection and reabsorption into the substrate.

Initial root zone pH was measured for 12 additional replicate containers without plants using the plug-squeeze method (Scoggins et al., 2002), during which leachate from each of the four cells per replicate was combined for measurements. Initial root zone pH was 5.84 ± 0.04 across the three solutions NH₄⁺/NO₃⁻ ratios. Final root zone pH was measured for each treatment replicate as described previously. Initial root zone pH (5.84) was subtracted from final root zone pH for each replicate to calculate change in pH units.

Change in pH was related to mEq of acidity or basicity produced in the root zone using a modified acid–base titration (Johnson et al., 2010), during which leachate from each of the four cells per replicate was combined for measurements. Initial root zone pH was 5.84 ± 0.04 across the three solutions NH₄⁺/NO₃⁻ ratios. Final root zone pH was measured for each treatment replicate as described previously. Initial root zone pH (5.84) was subtracted from final root zone pH for each replicate to calculate change in pH units.

Plant growth was measured as root and shoot dry weight gain during the experiment. Root and shoot tissue from seedlings and from final replicates was oven-dried for 48 h at 70 °C for dry weight determination. Dry weight gain was calculated by subtracting the initial dry weight of four seedlings from the total final dry weight for each replicate (four plants per replicate) of the same species, and was then divided by four to estimate dry weight gain per plant. For each replicate, species milliequivalents of acidity–basicity per plant was divided by the dry weight gain per plant to calculate mEq of acidity–basicity per gram dry weight gain for subsequent analysis.

The combined dry root and shoot tissue collected from each species replicate supplied with 0.100 solution was weighed and analyzed for nutrient concentration, to determine uptake of mEq of cations and anions. Differentiating between NH₄⁺-N and NO₃⁻-N uptake by tissue nutrient analysis was not possible without labeled nitrogen, and therefore cation/anion uptake was evaluated only with 0.100 solution, for which all N uptake was assumed to be in the anionic NO₃⁻-N form. Before oven-drying, roots were washed with phosphate-free detergent to remove substrate particles, and all tissue was rinsed in 0.1 N HCl followed by deionized water. Macronutrient and micronutrient concentrations were measured using inductively coupled plasma atomic emission spectrophotometry (Quality Analytical Laboratories, Panama City, FL). The weight of individual nutrients in plants was determined by multiplying plant dry weight by tissue nutrient concentrations. The total weight of each nutrient taken up per replicate during the experiment was determined by subtracting the total nutrient weight in four seedlings (initial) from the final nutrient weight for each replicate of the same species. For each replicate, uptake of nutrient weight values were divided by four to calculate uptake per plant, and were then divided by dry weight gain (measured in grams) per plant for mEq of nutrient uptake per gram weight gain.

Nutrient uptake was analyzed as mEq of net cations or anions taken up per gram dry weight gain, which was determined by subtracting mEq of net anions minus cations taken up per gram dry weight gain per plant for each replicate (resulting in negative values for a net uptake of cations or positive values for a net uptake of anions). Cation/
anion uptake was also evaluated as cation:anion uptake ratio, and was calculated by dividing mEq·g⁻¹ of net cations by mEq·g⁻¹ of net anions taken up. Nutrients taken up by roots were assumed to be NO₃⁻, H₂PO₄⁻, SO₄²⁻, MoO₄²⁻, Cl⁻ for anions and K⁺, Ca²⁺, Mg²⁺, Fe³⁺, Mn²⁺, H₂BO₄⁻, Cu²⁺, Zn²⁺, Na⁺, and Al³⁺ for cations.

Analysis of variance (ANOVA) with PROC GLM (SAS 9.4; SAS Institute, Cary, NC) was used to evaluate plant species and fertilizer NH₄⁺:NO₃⁻ ratio main and interaction effects on grams of dry weight gain per plant and mEq·g⁻¹ net acidity or basicity produced in the root zone. PROC GLM ANOVA was also used to evaluate species main effects on the cation:anion uptake ratio and mEq·g⁻¹ of net cation or anion uptake for plants supplied 0:100 NH₄⁺:NO₃⁻ solution. Mean separation used Tukey’s honestly significant difference (HSD) at the α = 0.05 significance level.

Hydroponic experiment. A factorial experiment with a randomized complete block design included a plant species factor with nine levels (arugula, basil, pepper, cucumber, eggplant, lettuce, spinach, tomato, and no plant control) and a nutrient solution NH₄⁺:NO₃⁻ N ratio factor with three levels (0:100, 15:85, and 30:70) using a modified 0.5x Hoagland’s solution at 7.14 mEq·L⁻¹. Species and cultivars were the same as mentioned previously in “Substrate experiment.”

On 27 Aug. 2015, 144-cell seedlings of arugula, basil, cucumber, eggplant, lettuce, spinach, and tomato (Knox Nursery, Apopka, FL) were transplanted into hydroponic culture vessels located in the same controlled-environment growth chamber with the same lighting conditions as in the substrate experiment. Daily air and solution temperatures during the hydroponics experiment were 24.1 ± 1.0 °C and 24.1 ± 0.9 °C, respectively.

Hydroponic culture vessels followed the design described by Dickson et al. (2016). Each hydroponic culture vessel was a 4.5-L white plastic container with a snap-on plastic lid. Substrate particles were washed from roots using deionized water before transferring seedlings to the culture vessels. Each seedling stem was wrapped with a black neoprene collar (diameter, 5 cm), which fit into a plastic mesh basket (diameter, 5 cm). Mesh baskets were supported in circular holes cut into the container lid, which allowed submergence of roots into nutrient solution held by the container. The neoprene collar reduced evaporation of the nutrient solution without constricting plant stems. A black plastic air tube inserted through each container lid aerated the nutrient solution constantly. Culture vessels were wrapped in aluminum foil to reduce light transmission to the nutrient solution, prevent algae growth, and help stabilize solution temperature. Each culture vessel initially contained five plants of either arugula, basil, cucumber, eggplant, lettuce, pepper, spinach, tomato, or no plants. Each culture vessel held 4.2 L nutrient solution and occupied a 25 × 25-cm area of growth chamber bench space.

Seedlings were initially supplied 0.25x modified Hoagland’s solution (3.57 mEq L⁻¹ N) with the 0:100 NH₄⁺:NO₃⁻ ratio for 7 d to acclimate to hydroponic conditions. After 7 d, the nutrient solution in each culture vessel was replenished with 4.2 L fresh 0.5x modified Hoagland’s solution at 7.14 mEq·L⁻¹ N (0:100 NH₄⁺:NO₃⁻), and culture vessels were replenished with fresh solution every 7 d thereafter. Solution pH was adjusted to 6.0 before transplant and for each fresh batch of nutrient solution using HCl and NaOH at 0.1 N. Solution pH was monitored every 2 d and maintained between 5.5 and 6.5. The nutrient solution was renewed every 2 d to acclimate plants to the different solution types.

On 16 Sept. 2015, culture vessels were replenished with 4.2 L 0.5x modified Hoagland’s nutrient solutions (7.14 mEq·L⁻¹ N) with NH₄⁺:NO₃⁻ ratios of 0:100, 15:85, and 30:70 for 2 d to acclimate plants to the different solution types. Modified 0.5x Hoagland’s solutions were mixed using reagent-grade salts and deionized water. Macronutrients were derived from (NH₄)₂SO₄, (NH₄)₆MoO₂·4H₂O, KH₂PO₄, MgSO₄·7H₂O, CaCl₂·2H₂O, KNO₃, and K₂SO₄. Micronutrient concentrations were constant at (μEq·L⁻¹) 72 Fe, 36 Mn, 6 Cu, 15 Zn, and 1 Mo derived from FeEDDHA (6.0% Fe, Akzo-Nobel); CuSO₄·5H₂O, MnSO₄·H₂O, (NH₄)₆MoO₃·2H₂O, and ZnSO₄·7H₂O. B exists as an uncharged molecule at pH less than 7 (Marschner, 2012) and was supplied at 1.0 mg·L⁻¹ from H₂BO₃.

The experiment started on 18 Sept. 2015 (day 0), and culture vessels were replenished with 4.2 L fresh 0.5x modified Hoagland’s nutrient solutions (adjusted to pH 6.0) with NH₄⁺:NO₃⁻ ratios of 0:100, 15:85, and 30:70. Species and solution NH₄⁺:NO₃⁻ ratio effects on solution acidity–basicity produced and individual nutrient uptake was measured for each replicate for 6 d. Each replicate consisted of one culture vessel. Each solution NH₄⁺:NO₃⁻ ratio and plant species combination (3 NH₄⁺:NO₃⁻ ratios × 9 species levels = 27 combinations) was replicated four times (27 combinations × 4 replicates = 108 total replicates). Two plants per replicate were harvested on day 0 for initial data collection, and the remaining three plants per replicate continued over the 6-d period and were harvested on 24 Sept. 2015 for final data collection.

Solution pH was monitored in each replicate every second day until final data collection on day 6, and pH was adjusted as described earlier. Initial and final solution samples were analyzed for individual nutrient ion concentration and were stored in the dark at 5 °C before analysis. Separate samples were collected for determining concentration of N forms. Nitrification was prevented in samples analyzed for N forms by adding sulfamic acid to lower the solution pH of samples to less than 3. Ammoniacal and NO₃⁻·N was measured by semiautomated and automated colorimetry (University of Florida Analytical Laboratory, Gainesville, FL). All other ions were measured by inductively coupled plasma atomic emission spectrophotometry (Quality Analytical Laboratories). Nutrient concentrations in solution were converted to mEq values for subsequent analysis.

For each replicate, plant growth was measured as grams of dry weight gain per plant during the 6-d period. At day 0, the two plants sampled destructively per replicate (culture vessel) were oven-dried at 70 °C for 48 h for initial root and shoot dry weight determination. The remaining three plants were sampled destructively on day 6 for final dry weight determination. For each replicate, the estimated initial dry weight per plant was subtracted from the final dry weight per plant to determine grams of dry weight gain per plant.

Net acidity or basicity produced in the root zone was calculated from the amount of acid (HCl) and base (NaOH) added per replicate during pH adjustment and the amount required on day 6 to titrate the solution back to pH 6.0. A positive net addition of acid (HCl) or base (NaOH) indicated that plants produced basicity and acidity, respectively. Titrated additions of acid and base were converted to mEq values. Net addition of acid or base was determined by subtracting the mEq of base added from the mEq of acid added per replicate or culture vessel, which was divided by three (three plants per replicate) and converted to mEq values to determine mEq of acidity or basicity produced per plant (a negative value indicated net acidity and a positive value indicated net basicity). For each replicate, mEq of acidity–basicity per plant was divided by the dry weight gain per plant and analyzed as mEq of net acidity–basicity per gram dry weight gain.

Individual nutrient uptake was calculated for each species–solution type combination by subtracting the mEq of each nutrient in solution for replicates containing plants from the average mEq in solution of the no-plant controls after 6 d. Cation vs. anion uptake was analyzed as mEq of anions taken up minus mEq of cations taken up for each replicate or culture vessel, where negative values indicated net cation uptake and positive values indicated net anion uptake. The assumed nutrient forms taken up were NH₄⁺, K⁺, Ca²⁺, Mg²⁺, Na⁺, NO₃⁻, H₂PO₄⁻, SO₄²⁻, Fe³⁺, Mn²⁺, Cu²⁺, Zn²⁺, MoO₄²⁻, Al³⁺, Cl⁻, and H₂BO₃⁻.

Net mEq cation/anion uptake per replicate was divided by three (three plants per replicate) for mEq of uptake per plant, then divided by the dry weight gain per plant (measured in grams) to determine mEq of net cation or anion uptake per gram dry weight gain. The cation/anion uptake ratio was calculated by dividing total mEq·g⁻¹ of cation uptake by total mEq·g⁻¹ of anion uptake.
Analysis of variance with PROC GLM (SAS 9.4, SAS Institute, Cary, NC) was used to evaluate plant species and NH₄⁺:NO₃⁻ ratio main and interaction effects on dry weight gain per plant, mEq g⁻¹ net acidity–basicity, mEq g⁻¹ net cation or anion uptake, and the cation:anion uptake ratio. Mean separation for ANOVA used Tukey’s HSD at the α = 0.05 significance level.

**Expected NH₄⁺:NO₃⁻ ratio for a neutral root zone pH.** Applied NH₄⁺:NO₃⁻ ratios for a neutral root zone pH were determined for each species in substrate and hydroponics (second objective), and analyzed as the percent NH₄⁻N of total applied N (remainder as NO₃⁻N) expected to result in a neutral pH. Linear regression was used to analyze the mEq g⁻¹ net acidity–basicity (dependent variable) at the three NH₄⁺:NO₃⁻ ratios (independent variable). In substrate, solutions 0:100, 20:80, and 40:60 corresponded to 0%, 20%, and 40% NH₄⁻N of total N, respectively. In hydroponics, solutions 0:100, 15:85, and 30:70 corresponded to 0%, 15%, and 30% NH₄⁻N of total N, respectively. The analysis was run by species, with separate analyses for substrate and hydroponic experiments. Each regression curve that was significant was then solved for percent NH₄⁻ of total applied N that would result in 0 mEq g⁻¹ net acidity–basicity (neutral pH).

### Results and Discussion

Plant growth differed between species in both experiments (P < 0.0001) and was affected by the NH₄⁺:NO₃⁻ ratio in substrate (P = 0.0128), but the interaction was not significant in either experiment (Table 1). In substrate, species ranged from 0.59 g dry weight gain per plant with arugula to 1.34 g dry weight gain per plant with eggplant. Dry weight gain was the least with the 0:100 solution (0.97 g) and greatest with the 40:60 solution (1.05 g). In hydroponic solution, species dry weight gain ranged from 0.65 g with spinach to 2.70 g with tomato (Table 1).

Plant species and solution NH₄⁺:NO₃⁻ ratio had significant main and interaction effects (P < 0.0001) on root zone acidity–basicity for plants grown in both substrate and hydroponic experiments, analyzed separately by experiment. In the substrate experiment (Fig. 1A), changes in substrate pH during the 33-d experimental period were converted to mEq of net acidity–basicity produced per gram dry weight gain per plant. Each species produced mEq g⁻¹ net basicity and increased pH when supplied with the 0:100 solution compared with other solutions, except for cucumber, which had no change in pH with 0:100. An increasing NH₄⁺:NO₃⁻ ratio increased the acidity produced by each species; the greatest mEq g⁻¹ net acidity in substrate occurred with the 40:60 solution. With the 40:60 solution, all species in substrate produced a net acidity and decreased pH except for arugula, which produced net basicity and increased pH in all solutions.

In the hydroponic experiment, species produced either net basicity (seven species) or had a neutral pH effect (spinach) with the 0:100 solution (Fig. 1B). Spinach also had a neutral effect with all solutions, indicating it was less sensitive to the NH₄⁺:NO₃⁻ ratio than other species in hydroponics. However, spinach had the lowest average dry weight gain in hydroponics (Table 1), and slight pH effects would be more difficult to measure experimentally. Similar to trends in Fig. 1A with substrate, arugula was the only species to produce basicity with 30:70 solution, and overall had the most basic effect on root zone pH of any species in each of the three nutrient solutions.

Characterizing species effects on root zone pH in units of mEq g⁻¹ net acidity–basicity allows comparison with research by Rengel (2003) and van Beusichem et al. (1988) for different species and NH₄⁺:NO₃⁻ ratios. Rengel (2003) compared 37 agronomic cereal and legume species by soil acidification, which ranged from 0.3 and 2.0 mEq g⁻¹ acidity (original units were reported as centimoles of H⁺ per kilogram shoot dry weight). Castor (Ricinus communis L.) grown as a model crop in hydroponic solution was shown to produce 3.70 mEq g⁻¹ acidity and 1.03 mEq g⁻¹ basicity when the nutrient solution contained either NH₄⁺-N or NO₃⁻-N as the sole N source, respectively (van Beusichem et al., 1988). Results from van Beusichem et al. (1988) are comparable to data in Fig. 1B. The 0:100 solution (100% NO₃⁻-N) resulted in a maximum of 1.31 mEq g⁻¹ basicity (for arugula) in hydroponics, and extrapolating from the 30:70 solution to 100% NH₄⁺-N (by multiplying by a factor of 3.33, calculated as 100% NH₄⁺-N + 30% NH₄⁺-N) resulted in up to 4.52 mEq g⁻¹ acidity for lettuce. These data may also be useful when predicting the mEq of acidity or basicity added into a substrate or hydroponic system when combined with a plant growth model.

The percent NH₄⁺-N of total applied N (remainder as NO₃⁻-N) expected to result in a neutral root zone pH for each species in either substrate or hydroponics was estimated using linear regression, and is summarized in Table 1. In substrate, cucumber produced a mEq g⁻¹ net acidity not different from zero when supplied with the 0:100 solution (Fig. 1A), which resulted in an estimated percent NH₄⁺-N value (2.3%) not different from 0.

### Table 1. Species gram of dry weight gain per plant, percent NH₄⁺-N of total supplied N expected to result in a neutral pH effect, and linear regression R² values analyzed separately for substrate and hydroponic experiments.

| Plant species | Substrate Dry weight gain (g)/plant | Hydroponic solution | Substrate Neutral NH₄⁺-N (%) | Hydroponic solution | Regression R² |
|---------------|-----------------------------------|---------------------|-----------------------------|---------------------|--------------|
|               |                                   |                     |                             |                     |              |
| Arugula       | 0.59 f                            | 0.87 c              | 46.0 ± 6.8                  | 23.3 ± 8.9          | 0.85         | 0.47         |
| Basil         | 1.09 cd                           | 1.44 bc             | 13.9 ± 4.1                  | 11.4 ± 5.5          | 0.81         | 0.64         |
| Cucumber      | 1.17 bc                           | 0.81 c              | 2.3 ± 5.5                   | 8.3 ± 5.4           | 0.83         | 0.75         |
| Eggplant      | 1.34 a                            | 2.06 ab             | 19.4 ± 5.0                  | 21.8 ± 10.1         | 0.81         | 0.72         |
| Lettuce       | 0.92 e                            | 1.34 c              | 12.5 ± 8.0                  | 6.6 ± 5.7           | 0.83         | 0.73         |
| Pepper        | 1.00 de                           | 1.25 c              | 26.5 ± 3.1                  | 14.9 ± 3.0          | 0.88         | 0.88         |
| Spinach       | 0.63 f                            | 0.65 c              | 25.9 ± 6.7                  | NA                  | 0.50         | NA           |
| Tomato        | 1.26 ab                           | 2.70 a              | 16.9 ± 3.7                  | 21.1 ± 5.5          | 0.83         | 0.71         |

NH₄⁺:NO₃⁻ ratio

- 0:100 (substrate and hydroponic) 0.97 b 1.44 a
- 20:80 (substrate) or 15:85 (hydroponic) 0.99 ab 1.46 a
- 40:60 (substrate) or 30:70 (hydroponic) 1.05 a 1.20 a

Species main effects

| NH₄⁺:NO₃⁻ main effects | ** | ** |
|------------------------|----|----|
| Species × NH₄⁺:NO₃⁻ interaction | NS | NS |

Dry weight gain data represent least-square means of 18 replicates per species and 48 replicates per NH₄⁺:NO₃⁻ ratio for the substrate experiment and least-square means of 12 replicates per species and 32 replicates per NH₄⁺:NO₃⁻ ratio for the hydroponic experiment. Mean separation for species and solution main effects on dry weight gain data used Tukey’s honestly significant difference at α = 0.05, and means sharing the same lowercase letter were not significantly different. Linear regression consisted of percent NH₄⁺-N of total N supplied (remainder as NO₃⁻-N) as the independent variable and milliequivalents of acidity–basicity per gram dry weight gain per plant as the dependent variable for each species. Error bars represent ±95% confidence intervals for percent NH₄⁺-N parameters. Linear regression slope coefficient used to calculate neutral percent NH₄⁺-N values was not significant for hydroponic spinach. * ** *** Not significant or significant at P < 0.05, 0.01, or 0.0001, respectively. NA = not applicable.
from zero (Table 1). The neutral percent NH₄⁺-N in substrate estimated for arugula (46.0%) was greater than the percent NH₄⁺-N in the 40:60 solution (40% NH₄⁺-N), which decreases the reliability of this estimate. In the case of cucumber in substrate, a complete 100% NO₃⁻-N fertilizer would therefore be needed for a neutral pH effect and to prevent a decrease in root zone pH. In contrast, other species were estimated to require a combination of NH₄⁺-N and NO₃⁻-N to achieve a neutral pH in substrate, where NH₄⁺-N ranged from 12.5% (lettuce) to 46.0% (arugula) of total N.

In hydroponics, the estimated neutral percent NH₄⁺-N ranged from 6.6% (lettuce) to 23.3% (arugula) of total N (Table 1). There was a positive relationship \( (r = 0.7 \text{ and } R^2 = 0.5; \text{data not shown}) \) between the neutral percent NH₄⁺-N estimated for hydroponics compared with the neutral estimated percent NH₄⁺-N in substrate when all species in Table 1 were included other than spinach (linear regression was not significant for spinach in hydroponics). This indicates that species such as arugula, which required a greater percent NH₄⁺-N relative to other species in substrate, would also require a greater percent NH₄⁺-N in hydroponics compared with a lower percent NH₄⁺-N for basil, cucumber, and lettuce. The estimated neutral percent NH₄⁺-N was intermediate for eggplant, pepper, and tomato.

For a neutral pH effect, recommended values for percent NH₄⁺-N of total N range from 5% to 10% in hydroponics and from 10% to 15% in substrate for most species (Sonneveld and Voogt, 2009). However, certain species have been reported to require percent NH₄⁺-N values more than or less than recommended ranges. For example, neutral percent NH₄⁺-N values of 0% and 25% were reported by Sonneveld and Voogt (2009) for melon and rose, respectively. Imas et al. (1997) found that tomato grown in nutrient solution required a neutral percent NH₄⁺-N of 30%, and 21.1% NH₄⁺-N was estimated for tomato in our hydroponic experiment (Table 1). Under similar experimental conditions with floriculture species, Johnson et al. (2013) found that petunia (Petunia ×hybrida Vilm.-Andr.), impatiens (Impatiens wallerana Hook. F.), and zonal geranium (Pelargonium xhortorum Bailey L.H.) required 31%, 10%, and 0% NH₄⁺-N, respectively, for a neutral pH in peat-based substrate. Based on our results, arugula was a particularly basic species compared with other species tested in this and other studies.

Substrates and nutrient solutions were formulated using hydrated limestone and zero-alkalinity water, respectively, which would result in low buffering against change in root zone pH. The NH₄⁺:NO₃⁻ ratio required for a neutral root zone pH is increased when irrigating with high bicarbonate concentration because of the basic and buffering effect on pH (Bailey, 1996; Fisher et al., 2014). The neutral NH₄⁺:NO₃⁻ ratio is decreased as total N concentration increases in the applied fertilizer solution because NH₄⁺ is a strong acid compared with NO₃⁻ as a base (Fisher et al., 2014). Substrate materials and cation exchange capacity are other factors that also contribute to acidity–basicty and buffering against changes in root zone pH (Marschner, 2012; Sonneveld and Voogt, 2009). A neutral NH₄⁺:NO₃⁻ ratio is therefore highly dependent on the plant species and other conditions, and quantifying pH acidic and basic factors would require a complex interacting model. In horticultural practice, this complexity necessitates regular pH monitoring and adjustment of fertilizer or acid–base correction because of the likelihood of pH drift over time.

Species grown in substrate and supplied the 0:100 solution (100% NO₃⁻-N) had greater uptake of anions than cations, based on tissue analysis, resulting in a cation/anion uptake ratio of less than one (Table 2). This result is consistent with previous reports of plants supplied with NO₃⁻-N only (Haynes, 1990; Kirkby and Knight, 1977; Marschner, 2012). Arugula had the lowest cation/anion ratio (0.54) and greatest net mEq of anions taken up per gram dry weight gain (2.16 mEq·g⁻¹ net anion uptake). A positive relationship was found between species mEq·g⁻¹ net anion uptake and mEq·g⁻¹ net basicity produced by roots (Fig. 2A). Therefore, species with greater uptake of anions than cations had a greater tendency to increase root zone pH.

In the hydroponic experiment, increasing the NH₄⁺:NO₃⁻ ratio increased the cation:anion uptake ratio and resulted in greater net uptake of cations [negative mEq·g⁻¹ net cation/anion uptake (Table 2)].

Species and NH₄⁺:NO₃⁻ ratio had main and interaction effects on the cation:anion uptake ratio \( (P < 0.01, \text{Table 2}) \). The significant interaction occurred because, in contrast to other species, in spinach the cation:anion uptake ratio did not increase with the NH₄⁺:NO₃⁻ ratio (data not shown).
As mentioned previously, spinach also had the least average dry weight gain and total nutrient uptake compared with the other species. Species and NH₄⁺:NO₃⁻ ratio (P < 0.05), but not their interaction, affected mEq·g⁻¹ net anion or cation uptake. Overall, arugula had the greatest mEq·g⁻¹ of net anion uptake (1.15) whereas lettuce had the greatest mEq·g⁻¹ of net cation uptake (0.42). Positive correlations (r > 0.8; data not shown) found between mEq·g⁻¹ of net cations or anions taken up and mEq·g⁻¹ of net solution acidity.

Table 2. Cation/anion uptake ratio and milliequivalents (mEq) net cation or anion uptake per gram dry weight gain per plant analyzed separately for substrate and hydroponic experiments.

| Expt. | Plant species | Cation:anion uptake ratio | mEq net cation or anion uptake/g dry wt gain/plant | Significance level |
|-------|---------------|---------------------------|-----------------------------------------------|-------------------|
| Substrate | Arugula       | 0.54 c                     | 2.16 a                                         | *** ***           |
|        | Basil         | 0.75 a                     | 0.65 bcd                                       |                   |
|        | Cucumber      | 0.79 a                     | 0.65 bcd                                       |                   |
|        | Eggplant      | 0.72 ab                    | 0.51 d                                        |                   |
|        | Lettuce       | 0.62 bc                    | 1.25 b                                         |                   |
|        | Pepper        | 0.71 ab                    | 0.73 bcd                                       |                   |
|        | Spinach       | 0.68 ab                    | 1.17 bc                                        |                   |
|        | Tomato        | 0.79 a                     | 0.56 cd                                        |                   |
| Significance level | **** | ***** | ***** | NS | NS |
| Hydroponic solution | Arugula       | 0.78 d                     | 1.15 a                                         | *** ***           |
|        | Basil         | 1.10 ab                    | -0.27 a                                        |                   |
|        | Cucumber      | 1.18 a                     | -0.24 a                                        |                   |
|        | Eggplant      | 0.94 bcd                   | 0.32 a                                         |                   |
|        | Lettuce       | 1.18 a                     | -0.42 a                                        |                   |
|        | Pepper        | 0.99 bc                    | -0.15 a                                        |                   |
|        | Spinach       | 0.90 cd                    | 0.88 a                                         |                   |
|        | Tomato        | 0.92 cd                    | 0.85 a                                         |                   |
| NH₄⁺:NO₃⁻ ratio | 0:100 | 0.77 b | 1.00 a | | *** |
|        | 15:85         | 1.10 a                     | -0.21 b                                        |                   |
|        | 30:70         | 1.12 a                     | -0.18 b                                        |                   |
| Species main effects | *** | * | *** | NS | NS |
| NH₄⁺:NO₃⁻ main effects | **** | *** | *** | NS | NS |
| Species × NH₄⁺:NO₃⁻ interaction | ** | *** | ** | NS | NS |

Substrate experiment data represent least-square means of six replicates per species supplied with a NH₄⁺:NO₃⁻ ratio of 0:100. Hydroponic experiment data represent least-square means of 12 replicates per species and 32 replicates per NH₄⁺:NO₃⁻ ratio. Mean separation for species and solution main effects used Tukey’s honestly significant difference at α = 0.05, and means sharing the same lowercase letter were not significantly different.

YCalculated as mEq of total anions minus mEq of total cations taken up per gram of dry weight gain per plant.

NS, *, **, *** Not significant or significant at P < 0.05, 0.01, or 0.0001, respectively

Fig. 2. Species milliequivalents (mEq) of net cation or anion uptake per gram dry weight gain per plant (x-axis) vs. mEq of acid (− value) or base (+ value) produced per gram dry weight gain per plant (y-axis). (A) Data from the substrate experiment with a NH₄⁺:NO₃⁻ ratio of 0:100, where symbols represent species least-square means from six replicates. (B–D) Data from the hydroponic solution experiment represent species least-square means from four replicates for each NH₄⁺:NO₃⁻ ratio.
and basicity produced for each of the three NH$_4^+$:NO$_3^-$ ratios indicated that species effects on root zone pH in hydroponic nutrient solution were largely the result of cation/anion balance (Fig. 2B-D).

In horticultural practice, commercial growers often produce multiple species in the same production space and it may not be practical to supply each species with a separate NH$_4^+$:NO$_3^-$ ratio for neutral pH. However, supplying species that differ significantly in their pH effect, such as cucumber and arugula, with the same fertilizer formulation is likely to result in pH drift and nutritional problems. In these scenarios, one option may be to group species by their general acidic or basic tendencies and then adjust the NH$_4^+$:NO$_3^-$ ratio for each group for an about-stable pH across species.

One limitation of the scope of our study was that experiments were conducted with young plants in their vegetative growth stage. Several authors have reported that stage of plant development influences uptake of individual cation and anion nutrients (Bugbee, 2004; Marschner, 2012; Sonneveld and Voogt, 2009), particularly for fruiting species such as tomato and cucumber. Cation/anion uptake and root zone acidity–basicity may change during the shift from vegetative to reproductive growth stage, which may affect NH$_4^+$:NO$_3^-$ ratios and neutral percent NH$_4^-$N values estimated for tomato, cucumber, eggplant, and pepper in this study.

The potential to adjust the NH$_4^+$:NO$_3^-$ ratio as a strategy to manage pH is limited because increasing NH$_4^-$N often inhibits the uptake of cations such as K$^+$, Ca$^{2+}$, and Mg$^{2+}$ (Bugbee, 2004; Lea-Cox et al., 1999; van Beusichem et al., 1988). Increasing NH$_4^-$N increases susceptibility to certain physiologic disorders related to poor Ca$^{2+}$ uptake and translocation within the plant, such as “tip burn” in lettuce and “blossom end rot” in tomato (Marschner, 2012; Sonneveld and Voogt, 2009). The influence of N form on Ca$^{2+}$ uptake was therefore evaluated in our study for species grown in hydroponic solution. Species differed in Ca$^{2+}$ uptake ($P < 0.0001$), and increasing the solution NH$_4^+$:NO$_3^-$ ratio decreased Ca$^{2+}$ uptake across all species ($P < 0.0001$), but there was no interaction ($P = 0.2692$). Figure 3A shows Ca$^{2+}$ uptake (in mEq of Ca$^{2+}$ per gram dry weight gain) was greatest with the 0:100 solution (1.13), intermediate with the 15:85 solution (0.90), and least with the 30:70 solution (0.78). Uptake of Ca$^{2+}$ was greatest for cucumber (1.54) and least for spinach (0.38) (Fig. 3B). In addition to limiting Ca uptake, the risk of ammonium toxicity increases as NH$_4^-$N increases greater than 20% of total N in hydroponic solutions (Sonneveld and Voogt, 2009). Based on the required neutral percent NH$_4^-$N values in Table 1, adjusting the NH$_4^+$:NO$_3^-$ ratio alone may not be an effective strategy to control pH for species such as arugula, eggplant, and tomato in hydroponics or arugula, pepper, and spinach grown in substrate, especially because nutrient solutions were formulated using zero-alkalinity water and substrate was incorporated with hydrated limestone to avoid high pH problems with certain species (Argo and Fisher, 2002).

**Conclusions**

This study clearly showed that both plant species and applied NH$_4^+$:NO$_3^-$ ratio influenced root zone pH in substrate and hydroponics, which has implications for nutrient management in commercial horticulture. Increasing the NH$_4^+$:NO$_3^-$ ratio resulted in increased acidity, lower pH, and increased cation uptake. Species differences in pH effects resulted from differences in cation/anion nutrient uptake, even when supplied with the same NH$_4^+$:NO$_3^-$ ratio. Adjusting the applied NH$_4^+$:NO$_3^-$ ratio is one potential strategy to balance species’ acidity–basicity for a neutral root zone pH. The percent NH$_4^-$N of total applied N expected to result in a neutral root zone pH was less than 10% for cucumber and more than 20% for arugula in both substrate and hydroponics, and overall these species were the most acidic and basic, respectively. The remaining species had neutral percent NH$_4^-$N values between 10% and 20% in substrate or hydroponics, and tended to be intermediate in their acidity–basicity. In substrate, cucumber produced acidity–basicity not statistically different from zero when supplied with the 0:100 solution, and was estimated to require a nearly 100% NO$_3^-$N fertilizer to prevent a decrease in root zone pH. On the other hand, arugula was especially basic and had an estimated neutral percent NH$_4^-$N of 46% in substrate. Plant species is only one parameter that influences root zone pH and the NH$_4^+$:NO$_3^-$ ratio. Other factors include irrigation water alkalinity, substrate components, nutrient concentrations, limestone type, and incorporation rate. The high level of pH change in our trials was partly the result of formulating substrates and solutions using hydrated...
limestone and zero-alkalinity water, respectively, which resulted in low buffering against change in pH. Although we showed that adjusting the NH4+:NO3− ratio can control pH, there are limits to this strategy because high NH4− can decrease the uptake of Ca2+ and other cations and promote ammonium toxicity, affecting plant quality negatively. Based on our results, adjusting the NH4+:NO3− ratio alone may not be an effective strategy to control pH for arugula, eggplant, and tomato in hydroponics or arugula, pepper, and spinach grown in substrate. Preventing pH drift in substrate and hydroponics requires regular pH monitoring in combination with adjusting NH4+:NO3− ratios, acid–base injection, and other pH management strategies.

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