Initial Substrate Moisture Content and Storage Temperature Affect Chemical Properties of Bagged Substrates Containing Poultry Litter Fertilizer

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Abstract. Bagged potting mixes can be stored for weeks or months before being used by consumers. Some bagged potting mixes are amended with organic fertilizers such as poultry litter (PL), although there is little knowledge about how these and other organic fertilizers release in the substrate while in storage. The objective of this research was to determine nutrient availability from an organic PL fertilizer in a bagged potting substrate stored at different temperatures and with varying initial moisture content (IMC). The base substrate composed of 60 sphagnum peat : 30 bark : 10 perlite (by vol.) amended with 5.5 g L⁻¹ dolomitic limestone and 0.5 g L⁻¹ granular wetting agent. This base substrate was either not amended with additional fertilizer [nonfertilized control (NFC)] or amended with a PL fertilizer (microSTART60, 3N–0.9P–2.5K) in its original pelletized form (PL-P) or ground (PL-G), or an uncoated prill fertilizer (UPF, 15N–6.5P–12.5K). Substrates had IMCs of 25%, 45%, or 65% (by weight) and were stored at either 20 or 40 °C. The UPF treatment resulted in lower pH, higher electrical conductivity (EC), and higher percent recovered nitrogen (N) compared with other treatments, as was expected with a readily soluble fertilizer. Poultry litter particle size had no effect on any of the measured chemical properties of the stored substrates. Both PL fertilizer treatments resulted in pH similar to or lower than the NFC. The two PL fertilizers had higher EC throughout the experiment (1.59–2.76 mS cm⁻¹) than NFC (0.13–0.35 mS cm⁻¹). Poultry litter fertilizer provided a stable source of N in bagged potting mix over a range of IMC and storage temperatures, with little change in total N released over time.

There is increasing demand from consumers and mass merchant retailers for products that are organic, more sustainable, or have lower carbon footprints. Consumer demand for organic gardening products is being driven primarily by increased demand from younger consumers for edible gardening and related products (Granderson, 2015). This demand includes bagged potting mixes. In the previously cited market study (Granderson, 2015), 30% of respondents who purchased organic gardening products specifically purchased “organic soil.” Mass merchants are also seeking to reduce their carbon footprint by demanding greater sustainability from their supply chain (Anonymous, 2017).

From the standpoint of their physical components, developing organic potting mixes is relatively easy, as most commonly used substrate components are eligible for organic labels, including peat, perlite, vermiculite, and coir (Wander, 2015). Providing an organic source of nutrition in bagged potting mixes is more difficult. Many organic nutrient sources are not immediately available for plant uptake. Organic fertilizers in substrates can come from many sources, although all can be categorized under three broad categories including plant-based fertilizers, animal-based sources, including manures, and some mined sources (Burnett et al., 2016).

Organic fertilizers have been used in soilless substrates and container production. Bi et al. (2010) showed that container-grown marigolds (Tagetes patula L.) with 1% to 2% (by vol.) broiler litter fertilizer had similar growth, flowering, and foliar nutrient concentrations as those grown with traditional controlled release fertilizer (CRF). Likewise, Mattson (2014) compared the production of several greenhouse crops in 10-cm pots using a traditional liquid feed program with three granular organic fertilizers preincorporated into the substrate and concluded that the organic fertilizers provided sufficient nutrients for these greenhouse crops and shortened the production periods. However, these studies and other authors (Fisher et al., 2016) warn that N mineralization from organic fertilizers is difficult to predict; and thus, managing fertility over time can be problematic.

Substrate moisture and temperature are two important factors affecting N availability from organic fertilizers. Griffin and Honeycutt (2000) used growing degree-day models to accurately predict N mineralization from poultry and other manure types in a sandy loam soil, demonstrating the importance of temperature on N mineralization rates. Malhi and McGill (1982) likewise showed reduced soil moisture and reduced nitrification in three different field soils; the highest rate of mineralization occurred with soil matric potential of –33 kPa and declined but did not stop at –1500 kPa. Numerous other studies have documented the effects of soil moisture and temperature on N mineralization in

Fig. 1. Moisture content (% by wt) of the bagged substrates stored in a growth chamber set to 20 (left) and 40 °C (right) during a 180-d storage period. Error bars represent the 95% confidence interval about the mean. IMC = initial moisture content.
soils, as reviewed by Agehara and Warncke (2005). However, virtually all of the N mineralization research has been conducted on field soils, where mineralization rates are also dependent on the inherent chemical and physical properties of each soil type (Chae and Tabatabai, 1986). Research on the mineralization of N from organic fertilizers in soilless substrates is lacking. Therefore, the objective of this research was to determine nutrient availability from an organic PL fertilizer in a bagged potting substrate stored at different temperatures and with varying IMC.

Materials and Methods

The base substrate was 60 sphagnum peat: 30 aged pine bark fines: 10 perlite (by vol.) amended with 5.5 g L\(^{-1}\) dolomitic limestone (ECOPHRST; National Lime and Stone Co., Findlay, OH) and 0.5 g L\(^{-1}\) granular wetting agent (AquaGro 2000G; Aquatrols, Paulsboro, NJ). The treatment design was a 3\( \times \)4 factorial arrangement with three moisture levels and four fertilizer treatments. Water was added to the base substrate to attain IMC of 25%, 45%, or 65%, calculated as the mass of water per mass of wet substrate.

The fertilizer treatments included the base substrate described previously with lime and wetting agent only as a NFC, a PL fertilizer (microSTART60, 3N–0.9P–2.5K; Perdue-AgriBusiness LLC., Seaford, DE) in its original pelleted form (PL-P) or ground (PL-G) or an (UPF, 15N–6.5P–12.5K; Yara, Oslo, Norway). The prills for the PL-P fertilizer were 99% smaller than 3.36 mm, whereas those in the PL-G were ground with a coffee grinder to be less than 1.19 mm. To investigate the differences in the nutrient availability between organic and inorganic fertilizers, UPF is added to compare with PL treatments. All fertilizer treatments were incorporated at 0.59 kg m\(^{-3}\) N.

After fertilizer treatments were mixed into the base substrate, 0.4 L of each was placed in 10 \(\times\) 15 cm, two mil sealable plastic bags (Royal Bag, Brooklyn, NY). Each bag was perforated with four holes to simulate the perforation of commercial bagged potting mixes. The bags were placed in a growth chamber (VWR Signature Diurnal Growth Chamber, Model 2015; VWR International, Radnor, PA) set to 20 or 40 °C. Because the growth chambers could not be replicated for practical reasons, differences in nutrient availability from the substrates between the two storage temperatures will not be compared statistically but will be discussed. There were four bagged replications of each fertilizer, temperature, and moisture level combination for each of five harvest dates at 2, 14, 30, 90, and 180 d after mixing.

Two days after the substrate treatments were mixed, but before they were placed in growth chambers for storage, four replication bags of each fertilizer and moisture treatment combination were analyzed for water extractable nutrients using a modified saturated media extraction procedure. Substrates were transferred each to a 1 L sealable plastic bag.
and 350 mL of reverse osmosis water was added. The substrate slurry was placed in a refrigerator and stored at 2 °C until the time of analysis.

At analysis, pH was determined with a pH/ion analyzer (MA 235; Metler Toledo, Columbus, OH) and EC with a conductivity meter (Fisher 06-662-61; Thermo Fisher Scientific). Samples were subsequently filtered through GF/F binder-free borosilicate glass fiber filter paper (Whatman Ltd., Kent, UK) to remove particles greater than 0.7 μm. The filtrate was poured into 5-mL autosampler vials, capped, and analyzed using ion chromatography (ICS 1600 Ion Chromatography System; Dionex, Bannockburn, IL) for concentrations of nitrate (NO₃⁻), ammonium (NH₄⁺), and phosphate (PO₄³⁻). A separate aliquot of each filtrate sample was filtered with blood serum filters (Porex Plastic-Evacuated 6211; Porex Corp., Fairburn, GA) with an effective filtration of 10 μm to determine total soluble N (TN, organic and inorganic forms) with a N analyzer (LECO TruSpec CHN 630; LECO Corp., Saint Joseph, MI). Three additional bags for each moisture level treatment were analyzed for moisture content by weighing the contents of the bag, drying in an oven at 110 °C, and placing under vacuum to facilitate extraction of the water from the substrate and through the filter paper. Filtrates were immediately placed in a refrigerator and stored at 2 °C until the time of analysis.

Data were subjected to analysis of variance and correlation analysis, when appropriate, using statistical software (SAS v9.3; SAS Institute, Cary, NC). Data were plotted as means with error bars representing the 95% confidence interval (CI) about the mean, using graphing software (SigmaPlot 12.5; SYSTAT Software, Inc., Chicago, IL).

**Results and Discussion**

Moisture content of bagged substrates was reported in a previous experiment conducted simultaneously with the one described herein (Jeong and Altland, 2017). Summarizing briefly from that previously reported data, moisture content of the bagged substrates changed over time (Fig. 1). There were significant interactions between IMC and day for bags stored in both 20 and 40 °C chambers (P < 0.0001, data not shown). Over the duration of the experiment, moisture in bags with 25%, 45%, and 65% IMC stored at 20 °C were reduced by 17%, 28%, and 13%, respectively, whereas those stored at 40 °C were reduced by 21%, 41%, and 60%, respectively.

Substrate pH was affected by the interaction between day and IMC as well as fertilizer type and IMC in the 20 °C environment, and the three-way interaction of the main effects in the 40 °C environment (Table 1). The fertilizer by IMC interaction in the 20 °C chamber was reflected primarily by the impact IMC had on substrate pH (Fig. 2). Substrate pH for both storage temperatures with 25% IMC ranged between pH 4.1 and 5.4 (with a few exceptions in NFC), whereas those with 45% IMC ranged between 4.9 and 6.5, and those with 65% IMC ranged between 5.2 and 7.0 (Fig. 2). Although there was a higher order three-way interaction between day, IMC, and fertilizer type in 40 °C chambers, pH response to IMC was similar to the 20 °C chamber, with the lowest pH in bags with 25% IMC. A similar response to IMC occurred in bagged substrates fertilized with a CRF (Jeong and Altland, 2017). Dissolution of dolomitic limestone added to potting mixes can cause gradual increase in media pH over time (Carll, 2004). The calcium and magnesium carbonate [CaMg(CO₃)₂] in dolomitic lime dissociates in water to form exchangeable Ca²⁺ and Mg²⁺, as well as carbon dioxide and additional water molecules (Lindsay, 1979). Although water is...
necessary for the reaction to occur, there is no literature addressing the impact of moisture level in soils or soilless substrates on lime reactivity. Although not compared statistically, substrate pH response to IMC followed a similar pattern in 20 and 40 °C growth chambers. Both PL fertilizer treatments resulted in pH similar to or lower than the NFC. Bi et al. (2010) likewise showed that two noncomposted PL-based organic fertilizers had similar or lower pH compared with NFCs in container-grown marigold. Poultry litter particle size had no effect on substrate pH, as means and 95% CI bars for both treatments (PL-P and PL-G) overlapped throughout the experiment in all six IMC and temperature conditions. The UPF treatment reduced pH throughout the experiment, and means were similar to or lower than all other treatments. Fertilizer effects on pH were likely due to the potential acidity of each product. A similar reaction occurred in a related study, in that substrates fertilized with a CRF had a similar or lower substrate pH compared with NFCs (Jeong and Altland, 2017). Although the fertilizers used in this study do not provide a labeled potential acidity or basicity, most agricultural fertilizers are acidic (Hershey, 1991; Jeong et al., 2016).

Day, IMC, and fertilizer type affected substrate EC in two-way interactions, but not three-way interactions, at both 20 and 40 °C (Table 1). EC of the NFC substrate ranged from 0.13 to 0.35 mS·cm⁻¹ (Fig. 3). The two PL fertilizers had higher EC throughout the experiment (1.59–2.76 mS·cm⁻¹) than NFC. Similar to pH, PL particle size had no effect on EC, as the 95% CI bars overlapped for the two treatments throughout the experiment. Electrical conductivities of PL-amended substrates were similar to those reported by Bi et al. (2010) when using similar amendment rates (0.518 kg m⁻³ N) of a different poultry fertilizer. The UPF treatment consistently resulted in higher EC (3.64–5.09 mS·cm⁻¹) than both PL treatments and the NFC. This indicates that the salts in UPF were quickly dissolved and readily available in soil solution, whereas salts from PL were released more slowly over time. Using the same brand of PL fertilizer, Mattson (2014) observed EC values of 0.6 mS·cm⁻¹ in PL-fertilized substrates, compared with an EC of 1.4 mS·cm⁻¹ in containers fertilized with an industry standard 21N–2.2P–8.3K water soluble fertilizer. Mattson (2014) goes on to warn that EC from organic fertilizers such as those based on PL can be difficult to interpret because water-soluble nutrients (salts) from these products become available more slowly over time; and therefore, low EC does not necessarily reflect low fertility.

Inorganic N (\(\text{NO}_3^- + \text{NH}_4^+\)) recovered in substrates was affected by the three-way interaction between day, fertilizer type, and IMC in both 20 and 40 °C (Table 1). Percent N recovery of PL and NFC treatments was at or near 0% throughout the experiment in all of the temperature and IMC treatment combinations (Fig. 4). Inorganic N recovery from UPF-treated substrates was higher than PL or NFC substrates. Although IN recovery from UPF-treated substrates was somewhat erratic over time and substrate conditions, all six of the IMC and temperature combinations resulted in >40% recovery by the conclusion of the experiment.

Total N was affected by the three-way interaction between IMC, fertilizer type, and day in both temperature conditions (Table 1). Percent TN recovered was at or near 0% for the NFC substrates, as expected (Fig. 5). The two PL treatments were again similar to each other throughout the experiment in all six IMC and temperature treatment combinations. Percent TN recovered from both PL treatments never exceeded 50%. Recovery from PL substrates was slightly higher with 25% IMC at both 20 and 40 °C. Percent TN recovered from PL substrates ranged from 10% to 20% with IMC of either 45% or 65%, whereas it ranged from 25% to 42% in substrates with 25% IMC. Recovered TN was highest in UPF treatments. Although these values fluctuated throughout the experiment, final %TN values ranged from 80% to 97% averaged across all IMC and temperature conditions. Percentage of recovered TN values greatly exceeded those for percent IN (Fig. 5 vs. Fig. 4), especially in UPF treatments. The differences in results between TN and IN data can be explained by the analytical methods and the mechanism of N immobilization in soilless substrates. After extracting the solution from the substrates using the modified SME procedure, an aliquot of the solution was designated for IN analysis using ion chromatography, whereas a separate aliquot was designated for TN analysis using ion chromatography.

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**Fig. 4.** The recovered water extractable inorganic nitrogen (\(\text{NO}_3^- + \text{NH}_4^+\)) from a substrate composed of 60 sphagnum peat : 30 bark : 10 perlite by volume with 5.5 g L⁻¹ dolomitic limestone. Data are expressed as mean percentage of the total mass of N applied via each fertilizer product. Substrates were either not amended with fertilizer [nonfertilized control (NFC)], or amended with 0.59 kg m⁻³ N of a poultry litter (PL) [microSTART60, 3N–0.9P–2.5K] that was either pelletized (PL-P) or ground (PL-G), or amended with an uncoated prilled fertilizer (UPF, 15N–6.5P–12.5K, Yara). Substrates had initial moisture contents (IMCs) of 65% (top), 45% (middle), or 25% (bottom) and were stored at either 20 °C (left) or 40 °C (right). Error bars represent the 95% confidence interval about the mean.
analysis using the CHN analyzer. Samples for IN analysis were filtered to 0.7 μm, which specifically detects and quantifies ionic compounds such as NO₃⁻, NH₄⁺, and organic nitrogen) from a substrate composed of 60 sphagnum peat : 30 bark : 10 perlite by volume with 5.5 g·L⁻¹ dolomitic limestone. Data are expressed as mean percentage of the total mass of N applied via each fertilizer product. Substrates were either not amended with fertilizer [nonfertilized control (NFC)], or amended with 0.59 kg·m⁻³ N of a poultry litter (PL) (microSTART60, 3N–0.9P–2.5K) that was either pelletized (PL-P) or ground (PL-G), or amended with uncoated prilled fertilizer (UPF, 15N–6.5P–12.5K; Yara). Substrates had initial moisture contents (IMCs) of 65% (top), 45% (middle), or 25% (bottom) and were stored at either 20 °C (left) or 40 °C (right). Error bars represent the 95% confidence interval about the mean.

Fig. 5. The recovered water extractable total nitrogen (NO₃⁻, NH₄⁺, and organic nitrogen) from a substrate composed of 60 sphagnum peat : 30 bark : 10 perlite by volume with 5.5 g·L⁻¹ dolomitic limestone. Substrates were either not amended with fertilizer [nonfertilized control (NFC)], or amended with 0.59 kg·m⁻³ N of a poultry litter (PL) (microSTART60, 3N–0.9P–2.5K) that was either pelletized (PL-P) or ground (PL-G), or amended with an uncoated prilled fertilizer (UPF, 15N–6.5P–12.5K; Yara). Substrates had initial moisture contents (IMCs) of 65% (top), 45% (middle), or 25% (bottom) and were stored at either 20 °C (left) or 40 °C (right). Error bars represent the 95% confidence interval about the mean.

In summary, there are several conclusions that can be made from using PL fertilizer in bagged potting mix. First, PL particle size had no effect on any of the measured chemical properties of the stored substrates. Particle size of the PL-P fertilizer used in this study was sufficiently small and uniformly distributed throughout the growing media to affect the nutrient content of the bagged potting mix without further particle size reduction.

Second, values for percent IN recovered (near zero) in the PL-fertilized substrates are not reflective of true plant-available N. Previous unpublished research with the same PL-fertilized potting mix used in this research showed that plants grew well compared with conventionally fertilized substrates, and thus would have access to some form of N (K. Jeong, personal observation). Nitrate and NH₄⁺ analyses with laboratory-grade analytical equipment or field-grade probes should not be used to evaluate the quality or characteristics of PL-fertilized potting mix. By contrast, EC and TN provided data more consistent with previous experience (K. Jeong, personal observation). EC values for PL-fertilized potting mixes ranged from 1.59 to 2.76 mS·cm⁻¹ throughout the experiment. Although there are no established standards for EC in bagged potting mix, an EC of 2.00 mS·cm⁻¹ is generally considered to be ideal (several commercial sources, personal communication). Percentage of TN recovered from the PL-fertilized potting mix ranged from 10% to 42% across treatments. Furthermore, these TN values remained relatively stable over time within an IMC and temperature treatment combination (Fig. 5). Unlike the nutrient load from a CRF-fertilized potting mix that increased steadily over time (Jeong and Altland, 2017), nutrient loads in PL-fertilized potting mix were more stable over the 180 d storage period.

Another practical application of this research is the potential use of filtering techniques to quantify N forms in soilless substrates. The different filtering techniques used in this research before the IC and CHN analyses were not deliberate but merely followed the standard protocols for each laboratory system. Our results suggest that filtration to include or exclude bacteria-immobilized N, as well as which analytical
data further show that the PL fertilizer used provided a stable source of N in bagged potting mix during storage. Our data show PL fertilizer incorporated into a bagged potting mix over a range of IMC and storage temperatures (20 and 40 °C), with little change in total N released over time. These findings contribute to understanding how organic or bacteria-immobilized N becomes available for plant uptake.

The objective of this research was to determine chemical changes (pH and EC) and nutrient availability from an organic PL fertilizer incorporated into a bagged potting mix during storage. Our data show PL fertilizer provided a stable source of N in bagged potting mix over a range of IMC and storage temperatures (20 and 40 °C), with little change in total N released over time. These findings contribute to understanding how organic or bacteria-immobilized N becomes available for plant uptake.

**Fig. 6.** The percentage of recovered water extractable phosphorus (P) from a substrate composed of 60 sphagnum peat : 30 bark : 10 perlite by volume with 5.5 g·L⁻¹ dolomitic limestone. Data are expressed as the mean percentage of the total mass of P applied in each fertilizer product. Substrates were either not amended with fertilizer (nonfertilized control [NFC]), or amended with 0.59 kg·m⁻³ of a poultry litter (PL) (microSTART®60, 3N–0.9P–2.5K) that was either pelletized (PL-P) or ground (PL-G), or amended with an uncoated prilled fertilizer (UPF, 15N–6.5P–12.5K; Yara). Substrates had initial moisture contents (IMCs) of 65% (top), 45% (middle), or 25% (bottom) and were stored at either 20°C (left) or 40 °C (right). Error bars represent a 95% confidence interval about the mean.

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