Establishment of Calibration Curves for Comparing Pour-through and Saturated Media Extract Nutrient Values

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Abstract. Most commercial and university substrate testing laboratories’ recommended floriculture nutritional values are based on the saturated media extract (SME) method. With the recent gain in popularity of pour-through nutritional monitoring, alternative recommended values are needed for nutrient analyses based on pour-through extracts. Pour-through nutritional values were compared to the SME values to develop calibration curves and recommended nutritional values. Euphorbia pulcherrima ‘Freedom Red’ Willd. ex Klotzch. were grown for two consecutive growing seasons in 16.5 cm plastic pots with Fafard 4 P root substrate and fertigated with 200, 300, or 400 mg·L⁻¹ N from a 13N–0.88P–10.8K fertilizer. Linear relationships existed and inverse calibration curves for pour-through and SME comparisons were developed for (r²): EC (0.98), NO₃⁻ (0.98), P (0.97 to 0.99), K (0.99), Ca (0.94 to 0.97), and Mg (0.91). In addition, recommended pour-through substrate value ranges were developed for comparison with SME values. The established calibration curves and pour-through substrate value ranges will allow substrate-testing laboratories to make nutritional recommendations based on pour-through extractions.

Pour-through was introduced as a quick and simple method to analyze potted-plant pH and electrical conductivity (EC) on-site (Wright, 1986; Wright et al., 1990; Yeager et al., 1983). Pour-through extraction occurs by the displacement of the bulk solution by distilled water poured over the top of a substrate as a means to obtain a nutrient sample. This method has been widely accepted by researchers and producers of woody ornamentals for approximately 20 years (Ruter, 1992; Wright, 1986; Yeager and Wright, 1981, 1982; Yeager et al., 1983). However, the greenhouse industry has been reluctant to accept this method until recent promotion at grower conferences and scientific meetings (D. Bailey, personal communication).

A possible reason for lack of acceptance may be that producers are already using a simple 2 water: 1 substrate (by volume) dilution slurry for on-site pH and EC testing. However, the 2:1 slurry does not generally provide a sample that is suitable for complete nutritional analysis due to the diluted nature of the substrate sample and inconvenience of shipping a slurry to a commercial or university laboratory. A pour-through extract has the advantage that it can be easily shipped to a commercial or university laboratory in a sealed container and since the substrate solution is unadulterated, it can be a preferred testing medium for plant available nutrients (Burd and Martin, 1923). The ability to use a single testing method for on-site pH and EC analysis and to provide a sample to commercial or university laboratories would be advantageous to producers. Advantages would include the nondestructive nature of the test, the simple sample collection, and a single set of nutritional standard values to monitor crop performance.

A limitation for pour-through acceptance has also been the lack of standard nutrient values to interpret proper nutrient concentrations in the substrate solution. Previous researchers have noted that pour-through or leachate samples and saturated media extract (SME) values, the current substrate analysis standard procedure used by many commercial or university laboratories (Dole and Wilkins, 1999; Lang, 1996), provide comparable results (Cabrera, 1998; Handreck, 1994; Hipp et al., 1979; Jarrell et al., 1979; Wright, 1986; Wright et al., 1990; Yeager et al., 1983). However, calibration curves that would allow the establishment of recommended pour-through values based on established SME values were not provided (Cabrera, 1998; Hipp et al., 1979; Jarrell et al., 1979; Wright, 1986; Wright et al., 1990; Yeager et al., 1983) or the methods used to obtain the sample were not representative of the procedures used by many greenhouse producers (Cavins et al., 2000; Handreck, 1994; Hipp et al., 1979; Jarrell et al., 1979).

Yeager et al. (1983) provided correlation coefficients based on comparisons of pour-through or SME extraction values to nitrogen, phosphorous, potassium, or dolomitic limestone applications. Calibration equations to compare the two testing methods directly were not provided. Similarly, Wright et al. (1990) presented correlations coefficients for pour-through or SME to tissue nutrient values, but did not compare the two testing methods directly. While several researchers have validated pour-through as an interpretable substrate test, calibration equations to compare the two testing methods directly are still needed. Calibration equations would allow commercial or university laboratories to make recommendations based on pour-through extracts from established SME standard values.

Substrates used in floriculture production are commonly peat moss or peat moss and pine bark based (K. Santner, personal communication). Yeager et al. (1983) data were based on 100% pine bark substrate without plants. Although Wright (1986), reported no difference in nutrient extraction between a 1 peat : 1 perlite (by volume) and pine bark (100%) substrate, nutrient retention may be differentially altered for each substrate by plant uptake. Lang and Elliot (1991) noted ammonium oxidation varied among soilless substrate types and that plants in the substrates affected oxidation. These considerations warranted additional investigation of pour-through.

Compton and Nelson (1997) noted that plants could drastically alter nutrient values in small substrate volumes and recommend testing 1 to 2 h after irrigation to obtain values representative of plant available nutrients. One to two hours allows for equilibration of the substrate solution. Standardized pour-through values based on conditions similar to those used by commercial floriculture producers that represent plant available nutrients are needed.

Wright et al. (1990) and Yeager et al. (1983) examined SME pH values obtained from the suctioned extract. Yet, Warrncke (1998) suggest the pH samples should be obtained directly in the slurry. Values for pH obtained in the slurry may differ from pH values obtained in the vacuumed extract. The vacuum extraction can change the CO₂ content of the solution, which may affect the pH (Lang, 1996; Van Leirop, 1990).

The objective of this study was to develop calibration curves of pour-through and SME nutrient values based on protocols used by commercial producers on containers with a common floriculture substrate and actively growing plants. Euphorbia pulcherrima ‘Freedom Red’ Willd. ex Klotzch. was chosen for this study as it is highest value potted flowering crop in the U.S. (U.S. Department of Agriculture, 2003). The values attained allow producers to use a single substrate solution extraction to achieve both on-site and complete nutritional analysis by a commercial or university laboratory. The calibration curves allow laboratories to develop standard pour-through nutrient values based on established SME standard values.

Materials and Methods

Euphorbia pulcherrima ‘Freedom Red’ Willd. ex Klotzch. rooted cuttings were transplanted on 15 Aug. into 16.5-cm plastic pots (one plant per pot) using Fafard 4P commercial...
substrate (Fafard Inc., Agawam, Mass.) and fertigated with 200, 300, or 400 mg·L⁻¹ N from a 13N-0.88P-10.8K Cal-Mag fertilizer (Scotts Co., Marysville, Ohio). Polyethylene covered greenhouses were set at 21/16 °C (day/night) and the plants were grown under natural days. Substrate samples were collected at 6 and 9 weeks (0 representing transplanting of rooted cuttings). Pour-through extractions, modified from Wright (1986) were made 1 h after irrigation using 75 mL of distilled water to displace about 50 mL of leachate (Cavins et al., 2000). Pour-through samples were tested for pH (Cole-Parmer lab pH/mv meter ± 0.01 pH units) and EC (Cole-Parmer lab conductivity meter ± 0.01 mS·cm⁻¹) immediately following extraction. Substrate samples were selected at the same time for modified SME (Warncke, 1998). About 350 cm³ of substrate from the middle of the container was saturated with distilled water until glistening, then allowed to equilibrate for 30 min. The pH was then measured in the slurry. After pH values were obtained, the samples were then vacuum filtered through a Buchner funnel using Whatman 40 filter paper and pH and EC were measured in the vacuumed extract. All extractions were preserved by refrigeration (5 °C) and the addition of 0.05 mL phenolmercuric acetate saturated solution until the completion of the study. Samples were then analyzed for NO₃⁻, NH₄⁺, and P photometrically (Cataldo et al., 1975; Chaney and Marbach, 1962; Murphy and Riley, 1962) and for K, Ca, and Mg using atomic absorption spectrophotometry (Christian and Feldman, 1970).

The study was a split plot design with a 2 × 3 × 2 × 2-factorial treatment combination. Two years served as the main plots, three nutrient levels, two extraction times, and two extraction methods were the subplots. There were five replications per treatment. Analysis of variance was obtained using procedure general linear models (α = 0.01) and the nutritional values were regressed and calibration equations developed using PROC REG (SAS Institute, Cary, N.C.).

Results and Discussion

The substrate pH did not differ between years or extraction procedures (data not presented). The lack of differences between years suggests that pour-through and SME are reliable, consistent methods for obtaining substrate pH. Surprisingly, there were no differences among extraction procedures (pour-through, SME slurry, and SME vacuum extract) for pH. Yeager et al. (1983) noted that pH values for pour-through and SME (suctioned) were similar. However, Lang (1996) noted that differences (0.1 to 0.5 pH units) are commonly obtained between SME pH values when they are taken from a slurry or vacuum extracted sample. These differences are attributed to partial pressure concentrations of CO₂ in the extract versus the atmosphere (Van Leirop, 1990). Evidently, our samples were allowed adequate time to equilibrate with atmospheric CO₂ after vacuum extraction to allow the pH values to return to the initial values prior to vacuum extraction.

As the fertilizer rate increased (200, 300, or 400 Fig. 1. Calibration curves and equations for comparison of pour-through and SME nutrient values for *Euphorbia pulcherrima* 'Freedom Red' for (A) EC, (B) NO₃⁻, (C) P—year 1, (D) P—year 2, (E) K, (F) Ca—year 1, (G) Ca—year 2, (H) Mg.
Increasing fertility rates predicted based on pour-through values. This calibration equation so that SME values can be values (Fig. 1A). Table 1 lists an inverse of the and a calibration curve was developed to predict values (Whipker and Hammer, 1997). The EC values between pour-through EC values (3.54 mg·L−1) from NO3− uptake (Raven, 1986) and carbonates (HCO3− and CO32−). As noted above, the fertilizer formulation used generally produces basic residues, which increases the substrate pH.

Electrical conductivity (EC) was not affected by year (data not presented). The lack of differences between years suggests that pour-through and SME are reliable, consistent methods for obtaining substrate EC.

The substrate EC was affected by extraction method. Pour-through EC values (3.54 mS·cm−1) were higher than the SME EC values (2.56 mS·cm−1). Pour-through EC values are generally higher due to less dilution compared to SME EC values (Yeager et al., 1983; Wright et al., 1990). The EC values between pour-through and SME extractions were compared and a calibration curve was developed to predict values (Fig. 1A). Table 1 lists an inverse of the calibration equation so that SME values can be predicted based on pour-through values. This inversion is necessary because laboratories will likely obtain pour-through nutrient values and want to predict the SME value. The predicted SME values can then be compared to established standard values.

Electrical conductivity (EC) was affected by fertilizer rate. Increasing fertility rates (200, 300, or 400 mg·L−1) increased EC values (2.09, 2.99, and 4.07 mS·cm−1, respectively). Similar trends have been noted by Yelanich et al. (1990) for pour-through.

Table 1. Inverse calibration equations for pour-through and SME comparisons. Pour-through extraction values are represented as the independent variable (x) and SME values are represented as the dependent variable (y).

| Nutritional parameter | Regression equation | Adjusted r² |
|-----------------------|---------------------|-------------|
| EC                    | Y = 0.74x – 0.05     | 0.98        |
| NO3−                  | Y = 0.71x – 25.13    | 0.98        |
| P                     | Y = 0.69x – 1.74     | 0.97        |
| Ca                    | Y = 0.63x – 23.39    | 0.99        |
| K                     | Y = 0.70x – 4.66     | 0.99        |
| Mg                    | Y = 0.59x + 1.27     | 0.91        |

*Equation is based on values obtained in year 1.
*Equation is based on values obtained in year 2.

mg·L−1), the pH decreased (6.12, 5.95, and 5.69, respectively). This pH decrease likely occurred due to the increased ammonium (NH4+) content of the fertigation solution; indicating that the uptake of NH4+ from the fertilizer, which acidifies the root environment (Marschner et al., 1991), was high enough to counteract the basicity effects of the high nitrate fertilizer formulation (155 kg of calcium carbonate [CaCO3] equivalent) (The Scotts Co., personal communication).

Sampling date affected pH such that the substrate pH increased from week 6 to week 9 (5.80 to 6.17, respectively) presumably due to build up of basic residues, such as hydroxide ions (OH−) from NO3− uptake (Raven, 1986) and carbonates (HCO3− and CO32−). As noted above, the fertilizer formulation used generally produces basic residues, which increases the substrate pH.

Similar trends have been noted by Yelanich et al. (1990) for pour-through. The EC values between pour-through EC values (3.54 mg·L−1) from NO3− uptake (Raven, 1986) and carbonates (HCO3− and CO32−). As noted above, the fertilizer formulation used generally produces basic residues, which increases the substrate pH.

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Electrical conductivity (EC) was affected by fertilizer rate. Increasing fertility rates (200, 300, or 400 mg·L−1) increased EC values (2.09, 2.99, and 4.07 mS·cm−1, respectively). Similar trends have been noted by Yelanich et al. (1990) for pour-through. The EC was not affected by sampling date (data not presented). During weeks 6 and 9 of production (0 representing potting day of rooted cuttings), plants are in a state of rapid vegetative growth (Whipker and Hammer, 1997). The lack of differences between sampling dates may be due to the plants actively consuming a majority of the supplied nutrients.

Similar to EC, NO3− values were not affected by year (data not presented), but were affected by extraction method. As with EC, pour-through NO3− values (478.8 mg·L−1) were higher than SME NO3− values (313.2 mg·L−1). The NO3− values of pour-through and SME extractions were compared and a calibration curve and inverse calibration equation were developed to predict values (Fig. 1C and D and Table 1, respectively). Table 1 lists an inverse of the calibration equations so that SME values could be predicted based on pour-through values.

Potassium extraction values were similar between years (data not presented). However, extraction method affected K values. Like EC, NO3−, and P, pour-through K values (336.8 mg·L−1) were higher than SME K values (295.9 mg·L−1). The lack of differences between sampling dates may be due to the plants actively consuming a majority of the supplied nutrients.

Table 2. Two-way interactions affecting P nutrient extraction values. Means are an average of 5 replications per treatment.

| Year | Week | P (mg·L−1) |
|------|------|------------|
| 1    | 6    | 15.80      |
| 2    | 6    | 15.51      |
|      | 9    | 189.01     |
|      | 9    | 141.67     |
| Interation |       | 0.0004     |

Table 3. Two-way interactions affecting P extraction values. Means are an average of 5 replications per treatment.

| Year | Rate (mg·L−1) | P (mg·L−1) |
|------|--------------|------------|
| 1    | 200          | 9.11       |
|      | 300          | 15.49      |
|      | 400          | 22.37      |
| 2    | 200          | 105.99     |
|      | 300          | 153.50     |
|      | 400          | 236.52     |
| Interaction |       | 0.0001     |

Table 4. Two-way interactions affecting Mg extraction values. Means are an average of 5 replications per treatment.

| Extraction | Rate (mg·L−1) | Mg (mg·L−1) |
|------------|--------------|-------------|
|            | 200          | 91.15       |
|            | 300          | 125.55      |
|            | 400          | 182.25      |
| SME        | 200          | 61.90       |
|            | 300          | 93.30       |
|            | 400          | 117.50      |
| Interaction | Extraction x rate | 0.0045 |

Table 5. Recommended pour-through substrate value ranges transformed from SME substrate values.

| Nutritional parameter | Pour-through | SME |
|-----------------------|--------------|-----|
| pH                    | 3.6–6.0      | 3.6–6.0 |
| EC (mS·cm−1)          | 2.8–4.8      | 2.0–3.5 |
| NO3− (mg·L−1)        | 180–320      | 100–199 |
| P (mg·L−1)           | 11–16        | 6–9   |
| K (mg·L−1)           | 220–360      | 150–249 |
| Ca2+ (mg·L−1)        | 330+         | 200+ |
| Mg (mg·L−1)         | 100+         | 70+   |

*Values adapted from Warncke and Krauskopf, 1983.
*pH values are equivalent for both extracts.
*Values based on year 1 data.
*Values based on year 1 data.
mg·L⁻¹) were higher than SME K values (232.2 mg·L⁻¹). A calibration curve and an inverse calibration equation were developed to predict values (Fig. 1E and Table 1).

Potassium concentrations increased with increasing fertilizer rates. Fertilizer rates of 200, 300, or 400 mg·L⁻¹ N produced substrate K values of 178.5, 280.4, and 394.6 mg·L⁻¹, respectively. Similar trends have been noted by Wright et al. (1990) for pour-through extractions.

As with EC and NO₃⁻, K substrate values were not affected by sampling date (data not presented). This is attributed to the poinsettias being in a state of rapid vegetative growth and consuming the same amount of K (relatively) during the 3-week interval (Whipker and Hammer, 1997).

Calcium substrate values varied between year 1 (348.1 mg·L⁻¹) and year 2 (140.6 mg·L⁻¹). Similar to P, the initial starter charge provided in the commercial mix caused the plants to mature, temperatures, irradiance, and humidity could have contributed to differences. Analysis conducted on the substrate prior to planting indicated the Ca values in year 1 were 277.0 mg·L⁻¹ (SME) versus 42.6 mg·L⁻¹ (SME) in year 2. This initial starter charge continued to affect Ca concentrations throughout the crop cycle. Argó and Biernbaum (1995) noted that containers irrigated only with tap water maintained near optimal levels of nutrients 42 d after initial irrigation.

Extraction method affected Ca values. Like EC, NO₃⁻, P, and K values, pour-through Ca values (295.4 mg·L⁻¹) were higher than SME Ca values (193.3 mg·L⁻¹) due to lack of dilution of the pour-through extract compared to SME extract (Yeager et al., 1983; Wright et al., 1990). The Ca values of pour-through and SME were compared and calibration curves and an inverse calibration equation were developed to predict values for year 1 and 2 (Fig. 1F and G and Table 1).

Fertilizer rate affected Ca concentrations. The 13N–0.88P–10.8K Cal–Mag fertilizer provides 6% calcium (12, 18, or 24 mg·L⁻¹ for the N rates used in this study). Therefore, as fertilizer rates increased (200, 300, or 400 mg·L⁻¹ N), so did the Ca substrate concentration (177.6, 237.4, and 318.1 mg·L⁻¹).

As with EC, NO₃⁻, P, and K values, Ca substrate values were not affected by sampling date (data not presented). This is attributed to the poinsettias being in a state of rapid vegetative growth and consuming the same amount of Ca (relatively) during the 3-week interval (Whipker and Hammer, 1997).

Magnesium (Mg) extraction values were not affected by year (data not presented). However, an extraction by rate interaction affected Mg substrate values (Table 4). Similar to EC, NO₃⁻, P, K, and Ca values, Mg pour-through values were higher than SME Mg values and the values increased with increasing fertilizer rates for both extraction methods. Pour-through Mg values were higher due to lack of dilution compared to SME Mg values (Yeager et al., 1983; Wright et al., 1990). The Mg values of pour-through and SME were compared and a calibration curve was developed to predict values (Fig. 1H). Table 1 lists an inverse of the calibration equation so that SME values could be predicted based on pour-through values. As with EC, NO₃⁻, P, K, and Ca values, Mg substrate values were not affected by sampling date (data not presented). This is attributed to the poinsettias being in a state of rapid vegetative growth and consuming the same amount of Mg (relatively) during the 3-week interval (Whipker and Hammer, 1997).

**Conclusion**

Calibration curves were established for EC, NO₃⁻, P, K, Ca, and Mg. Due to lack of differences in our study, pour-through pH versus SME pH, and pour-through pH versus SME were not developed.

Differences between years were noted with P and Ca. This was attributed to the initial starter charge difference in the commercial substrate. For P, year 1 had lower initial concentrations than year 2. This lower concentration likely allowed P to bind with more affinity to substrate particles due to a lower ionic strength substrate solution. Since P was bound to particles with more affinity, there was less P freely available to the unadulterated pour-through substrate solution. Therefore, pour-through P extraction values were not as high, relative to SME values, in year 1 compared to year 2. The year 1 calibration curve should be used for P comparisons, as most substrate levels will be <100 mg·L⁻¹ P. The P values presented in Table 5 were based on the calibration curve obtained in year 1.

For Ca, year 1 had higher initial concentrations than year 2. This higher concentration may have saturated the substrate binding sites. Therefore, when the substrate samples were diluted (SME), a greater amount of Ca was able to move into the bulk solution (year 1) than would have if the binding sites were not saturated (year 2). The result was that pour-through and SME Ca values were closer in year 1 than in year 2. The year 1 calibration curve should be used for Ca comparisons, as most Ca substrate values will generally be higher than 250 mg·L⁻¹. The Ca values presented in Table 5 were based on the calibration curve obtained in year 1.

In agreement with Yeager et al. (1983) and Whipker and Wright et al. (1990), a relationship exists among pour-through and SME values. In our study, pour-through nutrient values (excluding pH and NH₄⁺) were about 1.4 to 1.6 times higher than SME values, except for Ca (year 1) whose relationship was about 1.2 times higher (Fig. 1A–H). This is similar to Yeager et al. (1983) who reported pour-through values about 1.3, 1.4, and 1.2 times higher than SME values for N, P, and K, respectively; Yeager also noted that pH values were similar, which is agreement with our findings. Scoggins et al. (2002) noted comparable findings with the press extraction method, which is similar to pour-through such that an undiluted substrate solution is obtained for analysis, with EC, NO₃⁻, P, and V values that were 1.4, 1.8, and 1.7 times higher than SME.

However, we did not find a similar relationship of pour-through and SME values as described by Wright et al. (1990) who noted that pour-through values were nearly double that of SME values. These differences are likely due to variances in testing procedures (water content of the SME) that may have altered nutrient values.

Based on the calibration curves developed in our study and recommended SME values (Warncke and Krauskopf, 1983), recommended pour-through nutrient values for greenhouse substrates have been developed (Table 5). Additionally, calibration curves (Table 1) that allow substrate-testing laboratories to convert pour-through values to SME values and compare those values to established standard values have been presented.

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