An Evaluation of Two Seedling Phenotyping Protocols to Assess pH Adaptability in Deciduous Azalea (Rhododendron sect. Pentanthera G. Don)

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Abstract. Deciduous azaleas are an important element of residential and commercial landscapes in the United States after substantial trait improvements to increase their market appeal. Despite progress in breeding for ornamental characteristics and cold hardiness, intolerance to elevated pH and calcareous soils continues to limit their use in managed landscapes. Therefore, we assessed the utility of in vitro and greenhouse phenotyping approaches to evaluate and select for improved soil pH tolerance to increase the efficiency of breeding for this important trait. The research presented offers an example for implementing image-based phenotyping to expedite cultivar development in woody ornamental crops.

Deciduous azalea (Rhododendron sect. Pentanthera G. Don) cultivars and species are highly ornamental shrubs that are valued in northern landscapes for their colorful and prolific flowers. One important group of deciduous azalea cultivars is the Northern Lights series, which since 1979 has grown to 15 interspecific hybrids including the original ‘Northern Lights’, ‘Mandarin Lights’, and ‘Lemon Lights’ that bring vivid pink, orange, and yellow colors into U.S. Department of Agriculture (USDA) hardiness Zone 4 northern landscapes (Hokanson, 2010). With nearly 630,000 plants sold at a combined wholesale value of $1.02 million between 2014 and 2016, these and other deciduous azalea cultivars represent a reliable source of revenue for woody plant growers who cater to landscape markets in cold climates (L. Caton, Sidhu and Sons, personal communication). Given the series’ popularity, the University of Minnesota breeding program continues to expand the series, with the first true red ‘UMNAZ633’ Electric Lights™ Red and double-flowered ‘UMNAZ493’ Electric Lights™ Double Pink introductions bringing new ornamental characteristics to cold-hardy woody ornamental germplasm (Hokanson et al., 2015).

Progenitors of the cultivars mentioned previously primarily resulted from crosses between North American deciduous azalea species native to the southern and eastern United States with European deciduous azalea hybrids (Mollis Azalea, Exbury, and Knapp Hill hybrids) from the early 20th century (Moe and Pellett, 1986). Each of these native species contributes unique ornamental attributes to cultivars released. These include pink flowers from wild-collected Rhododendron prinophyllum (Small) Millais and large, durable, orange flowers from Rhododendron calendulaceum (Michx.) Torr. (Moe and Pellett, 1986; Widlrechner, 1982). Additional ornamental characteristics such as fragrance from Rhododendron atlanticum (Ashe) Rheder and white flowers from Rhododendron viscosum (L.) Torr. were also introgressed into the breeding germplasm (Hokanson, 2010). The combination of traits from each of these species and their segregation in seedling populations has enabled a diverse series of Rhododendron cultivars to be developed for northern areas of the United States, where cultivation of the genus was historically thought to be impossible because of extreme winter cold temperatures (Widlrechner, 1982).

The greatest systematic breeding effort has been focused on improving the cold hardness of floral and vegetative tissues (Moe and Pellett, 1986). Initial cold hardness evaluations required substantial screening time in the field as the initial parental germplasm and progeny evaluations that led to ‘Northern Lights’ spanned the years 1957–79, a 22-year period. These initial efforts resulted in a series of deciduous azalea cultivars that are reliably cold-hardy in USDA Zone 4 (Hokanson, 2010). As cultivation of deciduous azaleas has spread throughout the Midwestern United States, tolerance to high pH or calcareous soils (calcium carbonate, CaCO3) that predominate in the region has become the most pressing trait to improve. High-pH soils severely limit deciduous azalea growth, primarily by restricting micronutrient availability, such as iron, for uptake by roots (Brady and Weil, 2004). Bicarbonate ions present in calcareous soils raise soil pH and keep the soil buffered, resulting in slower acidification of soil, further limiting micronutrient availability (Brady and Weil, 2004). This reduction in iron uptake on high-pH or calcareous soils leads to chlorosis and decreased photosynthetic capacity, which over time reduces vigor and increases mortality of deciduous azaleas in the landscape (Galle, 1974). Improving adaptability to high-pH or calcareous soils (henceforth referred to as pH adaptability) of cultivars would reduce plant maintenance and open up new markets in continental climates characterized by calcareous or high-pH soils.

Although some systematic selection against iron deficiency chlorosis has occurred in evergreen Rhododendron, the variation is hard to phenotype consistently because of confounding nutrient deficiencies and measurement error using a qualitative (1–10) rating scale in breeding populations (Dunemann et al., 1999; Preil and Ebbinghaus, 1994). Furthermore, prospects for improving pH adaptability in deciduous azaleas are slowed by long generation times and limited timeframe (at leaf flush) for observing symptoms of high pH stress such as iron deficiency chlorosis. Like many woody ornamentals, deciduous azaleas take
between 8 and 10 months from crossing to seed germination, and an additional 2 years before plants reach maturity for field trials (Susko, 2016). Growing plants in containers before planting in the field requires space and labor that is often a limiting factor in determining how many seedlings can be produced for screening each season. Therefore, selection at the early seedling stage could both reduce the number of plants carried forward and expedite breeding deciduous azaleas.

Rather than waiting for the manifestation of chlorosis in whole plant experiments, characterization of the root–soil interface (rhizosphere), including localized reduction in pH (rhizosphere acidification), could more quickly indicate whether certain genotypes will successfully acquire micronutrients and mitigate symptoms of high pH stress (Guerinot and Yi, 1994). Known as Strategy 1 iron acquisition, dicot plants extrude protons into the rhizosphere to create favorable conditions for iron solubility and reduction that enable its translocation into root cells (Brady and Weil, 2004; Briat and Lobréaux, 1997). Direct acidification of the rhizosphere by proton extrusion by plasma membrane H+-ATPase is a well-documented response to iron deficiency in dicot plants (Yi and Guerinot, 1996). Indirect acidification of the rhizosphere through other nutrient exchanges such as nitrogen is also possible (Haynes, 1990). Specifically, ammonium uptake also occurs in concert with proton extrusion, and thus, results in acidification of the rhizosphere (Escobar et al., 2006). Regardless of the exact mechanism by which rhizosphere acidification is initiated, quantifying phenotypic variation in rhizosphere acidification capability could facilitate identification of selections capable of acquiring sufficient iron to promote healthy growth under calcareous soil conditions.

Given the quantitative nature of pH adaptability identified in other plants (Froechlich et al., 2016; Shi et al., 2013), the use of image-based phenotyping methods could improve our ability to more quickly and more precisely identify selectable genetic variation for traits such as rhizosphere acidification that play a role in pH adaptability. Image-based phenotyping protocols are potentially well suited toameliorating the problems (i.e., subjectivity, slow speed) inherent in phenotyping pH adaptability. Image-based screening approaches have been used in many other crops to improve quantification of challenging traits, including root architecture, water use efficiency, and nutrient deficiencies while maintaining yield goals (Berger et al., 2010; Clark et al., 2013; Shi et al., 2013). Systematic phenotyping protocols could then be used to identify superior genotypes within existing breeding germplasm or in wild populations to identify new sources of tolerance for elevated pH for deciduous azaleas.

To address the challenges of identifying pH adaptability in deciduous azaleas, we evaluated an in vitro phenotyping protocol to identify quantitative variation in pH change in the root rhizosphere. We focused our screening methods on in vitro plants in an effort to increase the efficiency of breeding elevated pH–tolerant plants. We tested seedlings from the same crosses in vitro using pH indicator dye to measure plant-induced changes to the culture medium pH in the presence of liming treatments. We also tested seedlings from the same families in the greenhouse to measure leaf area change in response to liming treatments. Leaf area was quantified using image analysis methods implemented in MATLAB (MathWorks, Inc., Natick, MA) and ImageJ (National Institutes of Health, Bethesda, MD) to detect any relationship between rhizosphere acidification in vitro and seedling size when progeny of the same family were grown in a greenhouse. Ultimately, we sought to identify seedling deciduous azaleas with improved pH adaptability for use as progenitors of future cultivars.

Materials and Methods

Germlaspm. Four maternal and two paternal parents were used to create seven full-sib families (Table 1). Based on pedigree records, the parents were not related through kinship. All of the parents used to develop the factorial design families were advanced, numbered selections from the University of Minnesota deciduous azalea breeding program, including progenitors and named selections in the Lights series (Hokanson, 2010; Hokanson et al., 2015). We harvested mature seed capsules approximately 5 months after crossing and subsequently stored them in a seed cooler at 4 °C for 6 months.

In vitro culture. Following capsule storage and dehiscence, all seed was surface sterilized for 30 min in a 2.5% Plant Preservative Mixture (PPM) (Plant Cell Technology, Inc., Washington, DC) solution using a magnetic stirrer. After rinsing seeds in distilled water, the seeds were plated on petri dishes (35 per plate) containing 20 mL of Woody Plant Medium with nutrients at half strength (McCown and Lloyd, 1981) (pH 5.7) and 10 µL of PPM sterilant. Germination occurred under fluorescent lighting (Sylvania 34W E34e bulbs; Oram Sylvania, Inc., Mississauga, CA) with a 24-h photoperiod at a room temperature of approximately 24°C. Following germination, the photoperiod duration of the fluorescent lighting was shortened to 16 h for 2 months as seedlings developed root systems and mature leaves. Screening for rhizosphere acidification began approximately 8–9 weeks following initial germination when seedlings were approximately 1 cm in height. The screening medium comprised half-strength Woody Plant Medium adjusted to pH 7.8 with sodium hydroxide and supplemented with 0, 310, or 610 mg·L⁻¹ CaCO₃ in the medium to assess the effect of lime concentration on rhizosphere pH changes.

To visualize pH change, the pH indicator phenolsulfophthalein (phenol red) was incorporated into the tissue culture medium at a final concentration of 90 µM before autoclaving. Ten milliliters of the previously described medium, including the phenol red, was poured into 100-mL Pyrex glass tubes. Individual plants from each family were transferred into these tubes that were then sealed with plastic caps and parafilm (Bemis NA, Neenah, WI). The tubes were organized in a completely randomized design under fluorescent lighting with a 16-h photoperiod throughout the screening process. The temperature of the culture room was maintained at approximately 24 °C throughout the germination and screening periods.

Greenhouse culture. Seeds from all families were sown in a greenhouse on sphagnum moss in 5” x 5” plastic flats under fluorescent lighting with an 18-h photoperiod and daily misting. Greenhouse temperatures were maintained at approximately 27 °C during the day and 21 °C at night. Once the seedlings developed mature leaves, they were fertilized once with a 200-ppm Peters Excel 21N–2.2P–16.4K (21–5–20) fertilizer solution (Grace-Sierra Co., Milipitas, CA). The seedlings grew in the plastic flats on sphagnum moss for 10 weeks, after which they were transferred to new plastic flats containing azalea growing medium (60% half-inch sieved pine bark and 40% peat) amended with 0%, 10%, or 20% powdered CaCO₃ by dry weight to achieve final medium pH levels of 5.7, 7.0, and 7.5. The growing medium was saturated with the Peters Excel solution mentioned previously on transplanting. The pH of the greenhouse medium was measured at the conclusion of the 4-week screening period. Medium pH for each of the treatments at the end of the screening period (5.7, 7.0, and 7.5) was determined by creating a 1:1 medium to water slurry that was stirred and then allowed to equilibrate for 15 min. The pH of the slurry was measured using a Mettler-Toledo Seven-Multi pH meter (Mettler-Toledo LLC, Columbus, OH) at the University of Minnesota Soil Testing Laboratory (St. Paul, MN).

Image analysis protocol (in vitro). Phenol red changes from red to yellow as the

Table 1. Seven families and the total number of seedlings per family grown in vitro for rhizosphere acidification screening and in the greenhouse for leaf area measurements.

| Paternal parent | Number of seedlings used for experiments (in vitro/greenhouse) |
|-----------------|---------------------------------------------------------------|
|                 | UMN AZ 223 | UMN AZ 376 |
| UMN AZ 180      | 37/171    | 43/99     |
| UMN AZ 181      | 32/90     | 45/171    |
| UMN AZ 493      | 39/45     | —         |
| UMN AZ 546      | 28/81     | 43/99     |

*No seed set.*
medium solution becomes more acidic between pH 7.5 and 5.5. To relate the hue measurements of the culture medium to approximate pH, we prepared a series of Pyrex tubes containing tissue culture medium of known pH containing the phenol red indicator that served as a standard scale. The Pyrex tubes containing progeny from the mating designs were photographed in a windowless room under fluorescent lighting (GE T8 Ultramax Eco XL 28W bulbs; General Electric, Fairfield, CT) using a Canon PowerShot G16 digital camera, with a 6.1–30.5 mm lens (Canon, Inc., Melville, NY), mounted on a tripod. Camera settings included an exposure time of 1/8 s, an International Standards Organization (ISO) speed of 80, a focal length of 6 mm, and a maximum aperture of 1.68.

Each culture tube in the experiment was photographed for analysis when the seedling was first transplanted and 3 weeks later, with the images saved in JPEG format. To assess changes in medium pH, the images were processed using an ImageJ macro that isolated and cropped the tissue culture medium picture to a 2 cm x 2 cm square image from the center of the tube (Susko, 2016a). A MATLAB (MathWorks, Inc., Natick, MA) script was used to quantify changes in the color of the phenol red pH indicator over the 3-week screening period (Susko, 2016b). Our script calculated the average hue, saturation, and value measurements for all pixels in the image. All hue, saturation, and value measurements were reported as decimal values ranging from 0.0 to 1.0, indicating percentage of degrees around the color wheel for hue, percentage of light for saturation, and percentage of darkness for the value. These values for each image were written to a .csv file by the MATLAB script (Susko, 2016b) for subsequent analyses. We present hue results in the context of the color wheel with a hue value of 0 representing pure red (pH value > 7.5) and numerical increases toward one indicating a change to orange (pH value 7.5–6.0) and yellow (pH value < 6.0). The change in hue represented change in pH, with an increase in hue indicating acidification of the medium.

**Image analysis protocol (greenhouse).** Photos were taken of seedlings after transplanting and 4 weeks later to assess the effects of the liming treatments on seedling leaf area. Images of seedling flats were taken using a Nikon D2X digital camera (Nikon Inc., Los Angeles, CA) mounted on a custom-built light box that maintained consistent lighting for uniform camera exposures. The light box was illuminated with four compact fluorescent bulbs (26 W; General Electric, Fairfield, CT). Manual camera settings (f-stop of f/4, exposure time of 1/200 s, an ISO speed of 200, focal length of 35 mm, and a maximum aperture of 1.6) were maintained for all pictures. Images were cropped using ImageJ macros to first reveal the seedling flat and then to divide each flat into 25 separate, uniformly sized sections so that the leaf area change of individual seedlings could be assessed (Fig. 1). Seedlings were distinguished from the medium by contrasting values and hues detected using a MATLAB script, which was initially optimized to detect seedling leaf tissue using the Image Thresholding tool within the MATLAB Image Processing Toolbox (Fig. 1) (Susko, 2016c). Specifically, only pixels with a value measurement between 0.36 and 1.0 and a hue measurement between 0.1 and 0.9, which represented green leaf surface, were kept for analysis, with all other pixels filtered out of the image because they either constituted greenhouse medium or severely discolored leaf tissue. The cropped images of individual seedlings were then analyzed in MATLAB to compute the leaf area of the seedlings (cm²) by scaling the resolution of the image against a known distance. We then computed the change in seedling leaf area by comparing the initial and week 4 images.

**Statistics.** All variance estimates were calculated using R version 3.3.2 (R Foundation for Statistical Computing, Vienna, Austria), using a linear model and analysis of variance (ANOVA) to compare genotypic and treatment effects among progeny. For the in vitro experiments, phenotypic means for medium hue change were averaged separately by treatment and then averaged by family across the three liming treatments and reported as a percentage of the color wheel (360°), with significance determined using a least significant difference test implemented using the R package Agricolae (De Mendiburu, 2014). We then regressed the change in leaf area for seedlings for each greenhouse liming treatment on seedlings grown in vitro and in the greenhouse by taking the average phenotypic value for each treatment in the greenhouse and in vitro experiments. This was performed to visualize the relationship between in vitro hue change and leaf area change in limed soil-grown plants and to measure the variation in seedling leaf area change accounted for by rhizosphere acidification.

Phenotypic and genetic variances for rhizosphere acidification were estimated using a factorial (partial diallel) crossing scheme. We estimated the narrow-sense heritability for rhizosphere acidification within mating designs as the proportion of progeny variance over the total genetic variance across the three liming treatments. We also estimated the significance of the maternal and paternal effect on the phenotype using a linear model with maternal and paternal genotype factors for hue change (rhizosphere acidification). All genetic variance estimations were made using the mean squares.
generated from the ANOVA on linear models by first estimating the full-sib progeny variance for the factorial design (Eq. 1) to calculate the narrow-sense heritability (Eq. 2) (Bernardo, 2010).

\[ V_d = 2V_{\text{progeny}} \]

\[ h^2 = 1 - \left( \frac{MS_{\text{progeny}}}{\text{MS}_{\text{progeny}} + \text{MS}_{\text{family}} + \text{MS}_{\text{Treatment}}} \right) \]

Results

In vitro rhizosphere acidification. Depending on the numbers available, between 28 and 57 seedlings per family were screened in vitro and 45–171 seedlings per family were screened in vitro and 45–171 seedlings per family were screened in the greenhouse (Table 1). No seed set was obtained from the cross between AZ 493 and AZ 376. We observed rapid germination and initial growth of in vitro seedlings, with most developing at least three mature leaves and branched root systems within 8 weeks after germination. On transferring to high-pH screening medium (pH 7.8) amended with CaCO₃ treatments, the initial hue values of the medium averaged 0.980. After 3 weeks, the average hue change was 0.029 across all families and liming treatments tested, indicating a general hue shift from red to orange, and thus a decrease (acidification) in medium (pH ≈ 6.8–7.0). Within each family, average hue change was consistently highest in the 0 mg·L⁻¹ CaCO₃ treatment (Table 2). The family UMN AZ 181 × UMN AZ 376 had the lowest observed hue change at 0.024 (pH ≈ 7.0), whereas the family UMN AZ 493 × UMN AZ 223 had the highest observed hue change at 0.050 (pH ≈ 6.6) (Table 2).

The effect of CaCO₃ was significant on observed hue change in vitro, with seedlings placed into the 305 and 610 mg·L⁻¹ CaCO₃ medium showing lower measured acidification of the medium (Table 3) (Fig. 2). Comparing family mean phenotypic values averaged across liming treatments revealed significant differences in hue change, with the family UMN AZ 493 × UMN AZ 223 showing a significantly higher level of hue change across liming treatments (P < 0.05) (Table 2). The maternal genotype effect on pH change was significant (P = 0.02), whereas the paternal genotype effect was not significant (P = 0.94) (Table 3). Using the progeny variance estimations from phenotypic values in Eqs. [1] and [2], we estimated a narrow-sense heritability (h²) across families for hue change of 0.38.

Greenhouse screening. Germination and initial growth of greenhouse-grown seedlings resulted in well-branched root systems and at least three mature leaves on each seedling before screening on liming treatments. Although progeny in each family differed slightly in foliage color before initiating high-pH/CaCO₃ treatments, the MATLAB image analysis scripts were successful in differentiating healthy, primarily green leaf tissue from the greenhouse medium background. This enabled us to quantify the area of green leaf tissue for each family at two time points. Mean initial green leaf areas were uniform for most families in the experiment, with only UMN AZ 180 × 223 and 181 × 376 having a significantly lower mean initial green leaf area (Fig. 3). Following the 4-week screening period, the effect of family on green leaf area change was highly significant across liming treatments (P < 0.001), with visible trends in seedling green leaf area apparent across families and liming treatments (Fig. 4). After 4 weeks, seedlings in the 10% and < 0.001), with visible trends in seedling green leaf area apparent across families and liming treatments (Fig. 4). After 4 weeks, seedlings in the 10% and 20% CaCO₃ by weight treatments were observed to have reduced leaf area and reddened or necrotic leaf tissue (P < 0.001) (Table 4; Fig. 1).

The 10% and 20% lime by weight treatments resulted in seedlings with lower and sometimes negative leaf area change (Fig. 4). This negative leaf area change indicates a reduction in the amount of green tissue over the screening period due to leaf death and/or severe discoloration which caused leaf hue to fall into the threshold of the image that was masked by the MATLAB analysis.

In comparing the average green leaf area change per treatment in the greenhouse with in vitro hue change in a given liming treatment for each family, in vitro hue change was positively correlated with green leaf area measured in the greenhouse (Fig. 5). In a linear model with in vitro hue change as the dependent variable, the calculated R² value was 0.45.

Discussion

This method of quantifying relative changes in rhizosphere pH may allow breeders of woody plants to more easily identify genotypes with superior capacity for iron uptake on high-pH or calcareous media. Although colorimetric visualization of changes in medium pH has been used in vitro for some time (Marschner et al., 1982; Yi and Guerinot, 1996), this experiment represents the phenotyping potential of such an approach in a high-throughput, quantitative breeding context. Although variation among families for rhizosphere acidification was significant, it is important to note the broad reduction in rhizosphere acidification across families growing in medium containing CaCO₃. This reduction in acidification in the presence of CaCO₃ is due to the buffering
CaCO₃ has long been documented (Hume, 1948) as increasing the solubility of micronutrients such as iron. Deciduous azalea intolerance to rhizosphere acidification is maternally inherited, an effect of bicarbonate ions on pH (Brady and Weil, 2004; Briat and Lobréaux, 1997). Thus, CaCO₃ limits total acidification of soil, and thus the ability of rhizosphere acidification to increase the solubility of micronutrients such as iron. Deciduous azalea intolerance to CaCO₃ has long been documented (Hume, 1948), and it is likely that the buffering of rhizosphere acidification contributes in part to the poor performance of deciduous azaleas on these soils. Nonetheless, quantifiable genetic variation for rhizosphere acidification and its positive relationship with seedling leaf area change suggest that genetic variation for alkalinity tolerance exists among deciduous azalea genotypes.

The narrow-sense heritability value of 0.38 for rhizosphere acidification we observed suggests that on average, if we were to select the seedlings from this population with the highest observed in vitro hue change and intermate them, we would expect our new population mean for rhizosphere acidification to increase by 38% compared with the previous population. However, the biological significance of the maternal and paternal components of the noted additive genetic variation should be interpreted cautiously, as a small number of parents (six) were used. It is possible that a more significant paternal genetic effect exists; however, it was not detected in this factorial design where only two paternal parents were used. Unless variation for rhizosphere acidification is maternally inherited, an unlikely scenario for a trait with such continuous variation (Bernardo, 2010), the inclusion of more paternal parents in a mating design will likely show a significant paternal effect on rhizosphere acidification variation.

Although increase in hue measured in vitro over time suggests that rhizosphere acidification occurs in deciduous azaleas, it does not confirm that it is a direct response to iron-limited conditions present in high-pH or calcareous soils. The rhizosphere acidification response in dicot plants (Strategy 1) is typically initiated in response to low iron in tissues and involves proton extrusion through reverse ATPase (Guerinot and Yi, 1994). Among woody genera, this direct rhizosphere acidification response has been confirmed in *Quercus* L., *Vitis* L., and *Prunus* L. (Gogorcena et al., 2001; Gonzalo et al., 2011; Ksouri et al., 2006), whereas it is absent in *Vaccinium* L. under iron-deficient culture medium in the absence of CaCO₃ (Nunez et al., 2015). The significant reduction in rhizosphere acidification in the presence of increased levels of CaCO₃ we observed across all deciduous azalea families was similar to that shown previously among *Vitis* L. cultivars (Ksouri et al., 2006). The observed acidification in our experiment could also have been indirectly caused by the uptake of ammonium cations, which were present in the Woody Plant Medium used. Active extrusion of protons is also a mechanism used by root cells to stabilize intracellular charge balances that facilitate cation uptake such as ammonium (Haynes, 1990). This subsequently acidifies the rhizosphere and occurs independently of proton extrusion due to iron deficiency (Escobar et al., 2006). An additional experiment looking at acidification in the absence of ammonium at high pH would be necessary to confirm a physiological basis for direct acidification of the rhizosphere under iron-limited conditions present in calcareous or high-pH media.

The reduced change in leaf area among greenhouse-grown seedlings in treatments with CaCO₃ relative to the treatment lacking CaCO₃ further demonstrates the effect of high pH stress induced by bicarbonate ions on deciduous azalea seedlings. In addition, the effect of family on greenhouse seedling leaf area change was also significant. This pattern of treatment and family differences in response to increased CaCO₃ mirrored that of the in vitro rhizosphere acidification experiment. Full-sib seedlings with high rhizosphere acidification response in dicot plants (Strategy 1) is typically initiated in response to low iron in tissues and involves proton extrusion through reverse ATPase (Guerinot and Yi, 1994). This subsequently acidifies the rhizosphere and occurs independently of proton extrusion due to iron deficiency (Escobar et al., 2006). An additional experiment looking at acidification in the absence of ammonium at high pH would be necessary to confirm a physiological basis for direct acidification of the rhizosphere under iron-limited conditions present in calcareous or high-pH media.

![Fig. 2. Boxplot depicting differences in hue resulting from CaCO₃ concentration in in vitro–grown seedlings.](Image)

Whereas there were significant differences between the control (0 mg·L⁻¹ CaCO₃) and the CaCO₃ treatments (*P* < 0.001), the difference in mean hue between the two CaCO₃ treatments was not significant.

![Fig. 3. Bar graph showing the original green leaf area mean and 95% confidence intervals for seedlings from each family used in the greenhouse phenotyping experiment. Unique letter combinations indicate a significant difference (*P* < 0.05) between family mean original seedling leaf areas.](Image)
of bicarbonate ions reduces seedling growth, and the variability in response among full-sib families suggests a genetic component to tolerance of high-pH, calcareous media.

Regardless of the exact mechanism by which we observed a pH change in the medium, the decrease in pH would nonetheless have consequences for iron solubility at the root–soil interface. The genotype specific variation, measurable narrow-sense heritability, and positive relationship between rhizosphere acidification and leaf area increase suggest that selection for in vitro hue change could serve as a means to improve pH adaptability in deciduous azalea. It is unknown at this point the degree of rhizosphere acidification that will occur in mature plants derived from this breeding population as mature plants will likely be subject to stronger genotype by environment interactions and genotype by year interactions that affect relevant phenotypes for pH adaptability. Determining mature plant response to elevated pH and high calcareous soils will require longer term observation of mature plants in field experiments. This could be accomplished via a multiyear observation of deciduous azalea growth rates under high pH stress and might shed light on the relationship between rhizosphere acidification in seedlings and the performance of genotypes in stressful landscape situations over time.

The high-throughput phenotyping of seedling leaf area change and rhizosphere acidification enabled data collection with relative speed in a nondestructive way, with methods readily transferable to future studies using the scripts presented here. Given market demand for a steady flow of new ornamental landscape cultivars, there is an incentive to accelerate the timeframes for evaluating plant germplasm. Speeding up the breeding process will require more efficient high-throughput phenotyping protocols (Van Nocker and Gardiner, 2014). Speed and accuracy, both of which will enable timely selection of azalea cultivars with improved pH adaptability, could be greatly improved through the use of image-based phenotyping (Furbank and Tester, 2011). The automated nature of image analysis through scripts removes potential ratings bias and saves substantial time. The scalability of image-based phenotyping to higher numbers of progeny also allows more data to be collected for less effort, which could lead to improved estimates of quantitative trait parameters and response to selection in breeding populations (Bernardo, 2010).

Broadly speaking, these methods should prove useful for quickly screening new germplasm, and when coupled with the use of a mating design, can estimate progeny variation for both rhizosphere acidification and seedling leaf area change. This will allow for estimations of quantitative trait parameters important for pH adaptability that were previously difficult to obtain for a woody plant. Such systematic application could be applied in Rhododendron to identify germplasm sources to develop durable cultivars for broader landscape use.

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**Table 4.** Analysis of variance (ANOVA) results for seedling leaf area change in the greenhouse liming experiment. Maternal and paternal effects are not separated in this ANOVA and are regarded together as the factor “Family.”

| Source of variation | df | Mean squares | F value | P value |
|---------------------|----|--------------|---------|---------|
| Family              | 6  | 1.004        | 9.031   | <0.001  |
| Treatment           | 2  | 0.883        | 7.948   | <0.001  |
| Family × treatment  | 12 | 0.903        | 8.127   | <0.001  |
| Residuals           | 949| 0.111        |         |         |

**Fig. 4.** Change in leaf area (cm²) of seedlings in seven families growing in three CaCO₃ treatments in the greenhouse over 4 weeks, measured through MATLAB image analysis. The effect of the liming treatments significantly reduced seedling leaf areas (P < 0.001).

**Fig. 5.** Relationship between in vitro average final hue for each treatment and family and greenhouse seedling green leaf area for each treatment and family. The dark shading around the regression line (R² = 0.45) denotes a 95% confidence interval. Individual dots represent a treatment average for each family, with treatments coded by color. The family UMNAZ 493 × 223 seedlings in the 10% and 20% CaCO₃ treatments, where average final hue explained significantly more than 45% of the variation in leaf area change on calcareous greenhouse media, are denoted by blue boxes.
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