RESEARCH PAPER

Evolution of US maize (Zea mays L.) root architectural and anatomical phenes over the past 100 years corresponds to increased tolerance of nitrogen stress

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Received 26 November 2014; Revised 29 December 2014; Accepted 4 February 2015

Abstract

Increasing the nitrogen use efficiency of maize is an important goal for food security and agricultural sustainability. In the past 100 years, maize breeding has focused on yield and above-ground phenes. Over this period, maize cultivation has changed from low fertilizer inputs and low population densities to intensive fertilization and dense populations. The authors hypothesized that through indirect selection the maize root system has evolved phenotypes suited to more intense competition for nitrogen. Sixteen maize varieties representing commercially successful lines over the past century were planted at two nitrogen levels and three planting densities. Root systems of the most recent material were 7° more shallow, had one less nodal root per whorl, had double the distance from nodal root emergence to lateral branching, and had 14% more metaxylem vessels, but total metaxylem vessel area remained unchanged because individual metaxylem vessels had 12% less area. Plasticity was also observed in cortical phenes such as aerenchyma, which increased at greater population densities. Simulation modelling with SimRoot demonstrated that even these relatively small changes in root architecture and anatomy could increase maize shoot growth by 16% in a high density and high nitrogen environment. The authors concluded that evolution of maize root phenotypes over the past century is consistent with increasing nitrogen use efficiency. Introgression of more contrasting root phene states into the germplasm of elite maize and determination of the functional utility of these phene states in multiple agronomic conditions could contribute to future yield gains.

Key words: Anatomy, architecture, density, maize, nitrogen, phenotype, root.

Introduction

Global food production must double in order to meet the demands of the future population of 9.6 billion people predicted by 2050 (Royal Society of London, 2009). There is a shortage of arable land (Pretty, 2008), so land use efficiency must increase dramatically to meet current and future demand. Maximizing crop nutrient and water use efficiencies is one approach for increasing land productivity (Lynch, 1998). Average maize production per hectare in the USA has increased ~8-fold in the past century (USDA National Agricultural Statistics Service, 2013). Genetic improvement and agronomic practices have contributed to this increased production about equally (Duvick, 2005). Increased yield in maize due to breeding has been associated with changes in a variety of phenes (i.e. basic units of phenotype, sensu...
Serebrovsky, 1925; Lynch, 2011; see York et al., 2013 for discussion), including phenology, leaf angle, and kernel number, among others (Tollenaar and Lee, 2006). Newer maize hybrids may also have increased tolerance to soil resource constraints such as low soil nitrogen (N) (McCullough et al., 1994a) and drought (Dwyer et al., 1992).

Over the past 100 years, maize breeding has continually adapted to changing agronomic conditions. Maize crop stands existed as widely spaced, open-pollinated varietal mixtures for thousands of years until the advent of hybrid seed encouraged genetic monocultures and the mechanization of farm tools necessitated spatial arrangements convenient for planting and harvesting between 1910 and 1940 (Bogue, 1982). During the same period, research on the role of plant nutrients encouraged farmers to amend soil with more N in organic and chemical forms (Russel and Williams, 1977). Subsequently, world chemical N use increased 3-fold between 1930 and the end of the Second World War, due mostly to application over a greater area (Frink et al., 1999). Since 1960 the use of chemical N fertilizer increased substantially in the USA from an average of 65 kg ha⁻¹ to 160 kg ha⁻¹, an increase of 240% (USDA Economic Research Service, 2014). In the early 1900s researchers were already aware that more productive land could accommodate a higher population density to optimize yield (Hume et al., 1908), and by mid-century the connection was made that N fertilization could support higher population densities to increase yield (Dungan et al., 1958). Maize population density increased from <30 000 plants ha⁻¹ in 1930 to 40 000 plants ha⁻¹ in 1960, and then doubling to an average of 80 000 plants ha⁻¹ in 2005 (Duvidk, 2005), and continues to increase. Maize in the USA has been selected to maximize yield in these changing intensive management systems, and evolution of the root system is hypothesized to have occurred to increase resource acquisition efficiency in these systems.

Maize performance is influenced by many root system architectural, anatomical, and morphological phenes and phene aggregates (see Lynch and Brown, 2012; York et al., 2013; Lynch et al., 2014) that influence root distribution and soil resource acquisition, including crown root number (York et al., 2013; Saengwilai et al., 2014b), topsoil foraging (Zhu et al., 2005a), crown root angle (Trachsel et al., 2013), rooting depth (Hund et al., 2008), lateral branching (Zhu and Lynch, 2004; Zhu et al., 2005b; Postma et al., 2014), root cortical aerenchyma (RCA) (Zhu et al., 2010a; Saengwilai et al., 2014a), living cortical area (Jaramillo et al., 2013), cortical cell size (Chimungu et al., 2014a), cortical cell file number (Chimungu et al., 2014b), and root hairs (Zhu et al., 2010b). Root phenes may also influence interplant competition and facilitation (Hodge et al., 1999; Ge et al., 2000; Bates and Lynch, 2001; Rubio et al., 2003; Postma and Lynch, 2012; Zhang et al., 2014). Interactions among root phenes for foraging utility and metabolism economics will determine how these root phenes influence soil resource acquisition together to create a functionally integrated phenotype (York et al., 2013). Simulation studies suggested that deeper roots which access more stored water may drive better performance of modern hybrids at higher plant densities (Hammer et al., 2009). An ideotype of maize root phenotypes for water and N acquisition has been proposed consisting of several interacting architectural, anatomical, and physiological phenes (i.e. steep, deep, and cheap, or ‘SCD’; Lynch, 2013). However, breeding in maize over the past 100 years has been primarily driven by selection for above-ground phenes and yield, with less understanding of root phene utilities and whether the root system has evolved over this period.

The authors hypothesized that breeders indirectly selected the maize root system over the past 100 years through the root system’s influence on yield. Specifically, the authors proposed that maize root architectural and anatomical phenes have evolved towards the states represented by the SCD ideotype (Lynch, 2013), which will be used here as a reference phenotype. The authors also hypothesized these root phenes exhibit plastic responses to varying N and density levels representing changes in US agronomic practices, but that plasticity will not disrupt the overall pattern of more contemporary maize cultivars exhibiting more SCD-like phenotypes. In order to test these hypotheses, 16 maize varieties spanning the past century were grown in two N levels and three densities.

Materials and methods

Plant material

One open-pollinated variety (OPV) and 15 hybrids (Table 1) were evaluated as representatives from the entire DuPont Pioneer Era panel (Duvick et al., 2004) which contains well-studied, commercially successful material from the past century in the USA. The release dates of the present material range from 1900 until 2011.

Experimental site

The experiment was conducted in four fields at the Russell Larson Research Farm (aka Rocksprings) of the Pennsylvania State University.

Table 1. Sixteen commercially successful varieties from the past 100 years in the USA were used in this experiment

| Era | Release | Breeding method |
|-----|---------|-----------------|
| 1   | Pre-1900| Open pollinated variety |
| 1   | 1934    | Three-way cross |
| 1   | 1936    | Four-way cross |
| 1   | 1941    | Four-way cross |
| 2   | 1946    | Four-way cross |
| 2   | 1952    | Four-way cross |
| 2   | 1953    | Four-way cross |
| 2   | 1954    | Four-way cross |
| 3   | 1982    | Single cross |
| 3   | 1991    | Single cross |
| 3   | 1999    | Single cross |
| 3   | 2002    | Single cross |
| 4   | 2006    | Single cross |
| 4   | 2008    | Single cross |
| 4   | 2008    | Single cross |
| 4   | 2011    | Single cross |
University (40°42’40.7”N, 77°57’11.1”W). The soil was a Hagerstown silt loam (fine, mixed, semi-active, mesic Typic Hapludalf).

**Treatment installation**

The experiment included four blocks, with each block being a separate 0.4 ha field. Each field was split in half randomly. On one half of each field, no N fertilizer had been applied since 2010 and had residual N of 13 mg kg⁻¹, while on the other half 145 kg ha⁻¹ of N was applied each year for the previous 2 years. In May 2013, entire fields received phosphorus (P) and potassium as determined by soil tests. On one half of each field, 145 kg ha⁻¹ elemental N as dry urea granules was broadcast to produce a high nitrogen (HN) plot, while the low nitrogen (LN) side received no N fertilizer. Subsequently, entire fields were disked to incorporate the fertilizers. A planter passed through all fields leaving behind non-planted rows in which to plant manually. Within these HN and LN split-plots, the factorial combinations of the three densities and 16 varieties were completely randomized. The three densities were 20 000, 40 000, and 80 000 plants ha⁻¹, which are abbreviated as 20K, 40K, and 80K throughout the manuscript and in the figures. Each N level, density level, and specific variety combination was planted in a five row plot that was 5 m long with a row distance of 76 cm. Seeds were hand-planted into rows marked by the planter using stakes and ropes marked to accommodate the various densities on 1 June 2013. Irrigation was not required because of adequate rainfall, insecticides were not required, and herbicide was applied before and after planting.

**Experimental sampling and harvest**

Shoots and root crowns were sampled from one block each day between 12 and 15 August 2013 (70 d after planting). Three entire plants were excavated and processed in the field from each plot using the shovelomics method (*Trachsel et al.*, 2011) with the shovel inserted 30 cm from the base of the plants. Shoots of the three plants were combined and partitioned into leaves, stalks, and immature ears, if any. The three root crowns were soaked in soapy water to disperse clay and then washed with a water hose and nozzle until most soil was removed. One root crown was selected for subsequent architectural measurements from the three root crowns that had been excavated based on its apparent average size and uniformity, and a second root crown was selected for anatomical sampling. All shoots were dried at 60 °C for 2 weeks and then weighed. Leaves were ground in a Wiley mill with a 40-mesh sieve (Thomas Scientific, Waltham, MA, USA) and a subsample analysed for N content with an elemental analyser (PerkinElmer 2400 Series II, Waltham, MA, USA) and a subsample analysed for N content with an elemental analyser (PerkinElmer 2400 Series II, Swedesboro, NJ, USA). Grain yield samples were collected from five random plants in each plot on 13 October 2013, threshed, and weighed.

**Root crown imaging and architectural measurement**

The original shovelomics method (*Trachsel et al.*, 2011) was modified to accelerate field processing while permitting more intensive measurements. Root crowns were kept in large plastic bins submerged in water inside a 5°C cold room until they were imaged within 1 week. Root crowns were imaged using digital cameras attached to frames with camera mounts from a height of 50 cm. Three identical cameras (Canon PowerShot A1200) operated by three researchers were used. Root crowns were placed under the camera on a matte black background. A 3 cm white plastic disk was included as a scale in every image, along with a printed sample label. Camera zoom and focus were kept locked for the duration of the imaging. An image was taken of the outermost layer of brace roots (above-ground nodal roots), and of the outermost layer of crown roots (below-ground nodal roots) by excising all whorls of brace roots. A representative nodal root of the respective whorl was excised from the side of the root crown and placed to the side of the root crown when imaged.

Image analysis was conducted in RSAJ (available at: [http://plantscience.psu.edu/roots/methods/computer/rsaJ](http://plantscience.psu.edu/roots/methods/computer/rsaJ)) which is a project for the ObjectJ plugin (available at: [https://sils.fnwi.uva.nl/bcb/objectj](https://sils.fnwi.uva.nl/bcb/objectj)) for ImageJ (*Schneider et al.*, 2012). RSAJ prompts the user to take sequential measurements from the images which are briefly described (see the RSAJ manual available at *JXB* online for elaboration). The same measurements were taken for the brace roots and the crown roots. Nodal root angle from the horizontal was derived trigonometrically from the stem width, maximum root crown width, and the height between stem width and root crown width. The number of nodal roots counted for a whorl was multiplied by two in order to account for the occluded half of the root system, based on previous observations. The diameter of the representative nodal root was measured at its base, along with the distance from where the representative root was excised from the shoot to where lateral roots emerge (distance to branching), which is equivalent to the distance from the stem to the first lateral. In order to calculate lateral root branching density, the number of lateral roots was counted along a measured length on the representative nodal root. Finally, the lengths of three representative lateral roots were measured and averaged for analysis.

**Anatomical sampling, imaging, and measurement**

One root crown per plot was processed for collecting anatomical samples. The middle part of a root from the second or third node that develops is the most representative for anatomical studies of maize roots (*Burton et al.*, 2013). Consequently, a nodal root from the second whorl was excised and a 4 cm segment was extracted 8 cm from the origin of the focal root from the stem and preserved in 75% ethanol. One root segment per sample was processed using laser ablation tomography (*Saengwilai et al.*, 2014a). Briefly, a sample is moved forward incrementally, the surface ablated or vaporized, and an image captured at the same time to generate a series of anatomical cross-section images. Three images for each segment were analysed with the semi-automated root anatomical measurement software RootScan (*Burton et al.*, 2012). See Table 2 for a list of and explanation of all reported architectural and anatomical root phenes.

**Table 2. Table of root phenes, their abbreviations (Abr), and their descriptions, divided into architectural and anatomical phenes**

| Architecture | Abr | Description |
|--------------|-----|-------------|
| Angle | Ang | Degrees from horizontal nodal root follows | |
| Number | # | Number of nodal roots in whorl | |
| Root diameter | Diam | Diameter of nodal root | |
| Distance to branching | DTB | The distance from cut to first lateral emergence on nodal root | |
| Lateral root density | LRD | Number of lateral roots within a centimetre of nodal root | |
| Lateral root length | LRL | Average length of three lateral roots of nodal root | |

**Anatomy**

**Cross-sectional area** | RXSA | The total cross-sectional area of a nodal root |
| Cortical cell area | CCA | The total area of a nodal root composed of cortical cells |
| Percent aerenchyma | %A | The percentage of the cortical area occupied by aerenchyma |
| Cell files | CF | The number of cell files within the cortex |
| Cell size | CS | The average area of individual cortical cells |
| Stele area | SA | The total area of the stele |
| Metaxylem vessel number | XV# | The total number of metaxylem vessels |
| Metaxylem area vessel⁻¹ | XVA | The average area of individual metaxylem vessels |
| Metaxylem area total | XVT | The total area of all metaxylem vessels |
Table 3. ANOVA table of shoot, crown root, and brace root architectural and anatomical measurements giving the F-value and significance for all factors and factor interactions

Architectural and anatomical phene abbreviations are as in Table 2, except amended with a crown root (CR) or a brace root (BR) prefix and including stem (St) width at that node. N level, density, and era are abbreviated N, D, and E, respectively, in the interaction terms.

| Factor       | N level | Density | Era | N:D | N:E | D:E | N:D:E |
|--------------|---------|---------|-----|-----|-----|-----|-------|
| DF           | 1       | 2       | 3   | 2   | 3   | 6   | 6     |
| Shoot mass   | 10.97*  | 505.52**| 1.47 NS | 1.04 NS | 2.11 NS | 1.35 NS | 0.79 NS |
| % Leaf N     | 379.46**| 54.61** | 2.03 NS | 5.37** | 0.62 NS | 0.36 NS | 1.83 NS |
| Total leaf N | 364.43**| 194.84**| 2.78*  | 9.56** | 0.93 NS | 1.04 NS | 2.46*  |
| Yield plant-1 | 110.06**| 451.89**| 52.55**| 6.33** | 3.07*  | 0.64 NS | 2.06 NS |
| Yield ha-1    | 142.84**| 147.46**| 68.70**| 74.86**| 7.66** | 7.95** | 4.13** |
| CR St width  | 2.68 NS  | 23.77** | 7.03** | 1.14 NS | 2.51 NS | 0.75 NS | 0.65 NS |
| CR #         | 0.12 NS  | 2.35 NS  | 1.77 NS | 0.54 NS | 1.34 NS | 1.15 NS | 1.55 NS |
| CR Diam      | 0.25 NS  | 4.37**  | 3.48*  | 0.08 NS | 0.55 NS | 0.27 NS | 0.89 NS |
| CR DTB       | 2.61 NS  | 2.61 NS  | 5.08** | 0.04 NS | 1.24 NS | 1.46 NS | 1.66 NS |
| CR LRD       | 1.46 NS  | 0.16 NS  | 1.28 NS | 1.21 NS | 0.73 NS | 0.48 NS | 0.63 NS |
| CR LRL       | 0.83 NS  | 3.46*   | 8.15** | 1.11 NS | 0.92 NS | 1.33 NS | 1.11 NS |
| CR Ang       | 0.30 NS  | 0.59 NS  | 9.13** | 1.12 NS | 0.26 NS | 0.48 NS | 0.98 NS |
| BR St width  | 29.71*  | 266.87**| 4.41** | 2.25 NS | 1.55 NS | 0.18 NS | 0.36 NS |
| BR #         | 37.45** | 41.54** | 2.10 NS | 2.02 NS | 1.36 NS | 0.45 NS | 1.16 NS |
| BR Diam      | 16.31*  | 43.15** | 0.90 NS | 1.08 NS | 0.23 NS | 0.58 NS | 1.24 NS |
| BR DTB       | 28.92*  | 61.96** | 6.13** | 1.12 NS | 2.41 NS | 0.29 NS | 1.51 NS |
| BR LD        | 3.71 NS  | 3.16*   | 0.08 NS | 0.76 NS | 0.12 NS | 1.24 NS | 0.80 NS |
| BR LL        | 3.1 NS   | 4.55*   | 1.41 NS | 1.32 NS | 1.57 NS | 0.54 NS | 0.61 NS |
| BR Ang       | 2.41 NS  | 4.12*   | 9.90** | 0.86 NS | 0.27 NS | 2.70*  | 0.69 NS |
| RXSA         | 1.34 NS  | 1.05 NS  | 1.81 NS | 0.1 NS | 0.91 NS | 0.32 NS | 0.2 NS  |
| CCA          | 19.13*  | 7.1**   | 2.92*  | 0.34 NS | 0.13 NS | 0.88 NS | 0.67 NS |
| %A           | 71.93** | 6.07**  | 1.71 NS | 1.67 NS | 0.27 NS | 1.2 NS  | 0.49 NS |
| CF           | 0.79 NS  | 1.56 NS  | 1.98 NS | 0.06 NS | 0.27 NS | 0.94 NS | 0.46 NS |
| CS           | 2.8 NS   | 1.64 NS  | 3.23*  | 0.14 NS | 0.18 NS | 1.19 NS | 0.63 NS |
| SA           | 0.94 NS  | 0.27 NS  | 1.68 NS | 0.06 NS | 0.48 NS | 0.52 NS | 0.26 NS |
| XVA          | 6.29 NS  | 2 NS    | 6.4**  | 0.74 NS | 0.22 NS | 1.76 NS | 0.84 NS |
| XVt          | 0.53 NS  | 1.68 NS  | 9.32** | 0.91 NS | 0.83 NS | 0.45 NS | 0.78 NS |
| XVT          | 3.94 NS  | 1.77 NS  | 0.53 NS | 0.06 NS | 0.37 NS | 0.57 NS | 0.86 NS |

*P<0.01; *0.01<P<0.05; NS P>0.05, not significant.

Simulation modelling

In order to investigate the relationship between changes in root system architecture over time and the functional utility for N acquisition, root systems representing the oldest and newest germplasm measured in this study were modelled in SimRoot (Lynch et al., 1997; Postma and Lynch, 2011a, b) growing in environments representing US fields a century ago and currently. SimRoot simulations include both a starting seed and soil conditions, where the soil is defined by soil, water, and nitrate properties. At the starting time, the seed produces root axes based on growth of real plants and with properties defined by the input files. In this study, all plant properties remained the same among simulations except for architectural and anatomical parameters as described below. The model includes a non-spatially explicit shoot model with expansion of leaf area leading to increased photosynthesis. Maximum growth rate is decreased proportionally to increasing nitrate stress, and nitrate stress will also increase the relative carbon allocation to the root system. The soil model SWMS_3D (Simunek et al., 1995) is used to simulate water and solute movement in the soil. The soil used in the current simulation was the same soil in which the field experiment took place with similar rainfall. The simulation time was 40 d.

An Old and a Modern phenotype were simulated which were identical except that the Modern phenotype was 10° more shallow angled and had one less nodal root in every whorl, which captures the essence of architectural changes observed in this study (Fig. 1). Three variants of the Old phenotype were simulated: (i) typical (Old); (ii) typical with a more shallow angle (Old+Angle); and (iii) typical with fewer nodal root numbers (Old+NRN). In this way, the contribution of each phene was assessed in isolation and combined. Two versions of the Modern phenotype were simulated: (i) typical (Modern); and (ii) typical with 40% RCA (Modern+RCA). Though RCA did not change among Era periods in this study, RCA might be an important phene for breeding programmes so was simulated at the maximum observed value in maize (Burton et al., 2013). These five phenotypes were simulated at a high density with HN availability (Current Environment), and at a low density with LN availability (Historic Environment), similar to the extremes of the field study.

Statistical analysis

In order to simplify data interpretation, the 16 varieties were grouped into four groups of four varieties (Era periods) based on their similarities of release year, original agronomic context, and breeding method (Table 1). All statistics were conducted and data graphics were created with R version 3.0.2 (R Core Team, 2013). Analysis of variance (ANOVA) was conducted with the mixed modelling package nlme (Pinheiro et al., 2013) in R with N level nested in a block as the random effect and with Era period, density, and N level as fixed effects. Post-hoc mean comparisons were conducted between Era periods when effects of release year, original agronomic context, and breeding method were significant (P<0.05) with Tukey HSD using the multcomp package (Hothorn et al., 2008). Principal component analysis (PCA) was conducted on complete observations of crown root architectural and anatomical raw data (n=261). Brace root architectural data
were excluded for PCA because many brace roots did not form lateral roots which substantially reduced the number of complete observations. Graphs were created in R with the ggplot2 package (Wickham, 2009).

Results

Shoot mass, nitrogen content, and yield

Total biomass produced on a per area basis did not differ among Era periods ($P=0.6122$, Supplementary Fig. S1 at JXB online), but plants grown under LN were 16% less massive than those under HN ($P=0.035$), and there was a 50.3% reduction in shoot mass per area from 80K to 20K ($P<0.0001$) (see Table 3 for complete ANOVA table for all results). The percentage N in the leaves was not affected by Era (Fig. 2A) but decreased 49.6% from HN to LN ($P=0.0003$), and increased 29.4% from 80K to 20K ($P<0.0001$). Grain yield on a per area basis (Fig. 2B) was significantly affected by all treatments, with a 52% reduction in yield from HN to LN ($P=0.0013$), a 66% increase from 20K to 80K plants ha$^{-1}$ ($P<0.0001$), and a 58% increase in yield from the oldest to the newest lines ($P<0.0001$).

Root system architecture

Brace root angle (Fig. 3A) decreased 12% from 59º to 52º from the horizontal ($P<0.0001$) from the oldest to the newest Era period, indicating that varieties in the most recent Era period are relatively more shallow angled than in the oldest. Brace root angle was more shallow angled at lower density, decreasing 5% from 57º to 54º from the horizontal between 80K and 20K ($P=0.0171$). Crown root angle (Fig. 3B) of the most recent material also decreased 10% from 62º to 55º from the horizontal, compared with the older materials ($P<0.0001$). A t-test demonstrates that crown roots are steeper than brace roots overall, with means of 58.4º and 55.6º from the horizontal, respectively ($P<0.0001$). Brace and crown root angles are >45º from the horizontal ($P<0.0001$). Distance to branching of both brace roots and crown roots increased considerably from older material to newer material (Fig. 3). For brace roots (Fig. 3C), distance to
branching increased 50% from 3.45 cm to 5.17 cm between the oldest Era period and the most recent ($P$=0.0005). Distance to branching was decreased from 6.16 cm to 2.7 cm from 20K to 80K planting density ($P$<0.0001), and decreased from 5.1 cm to 3.65 cm from HN to LN ($P$=0.0126), in brace roots. For crown roots (Fig. 3D), distance to branching increased 57% from .73 cm to 1.15 cm between the oldest Era period and the most recent ($P$=0.0019), but was not affected by density or N level.

Lateral root length was measured for three random lateral roots on each excised representative root and those three values averaged for subsequent analysis. For brace roots (Fig. 3E), lateral root length was not affected by Era period or N level, but was affected by density ($P$=0.0117), decreasing from 3.6 cm to 2.7 cm from 20K to 80K. However, the length of crown root lateral roots (Fig. 3F) was significantly affected by Era period, increasing 29% from 3.73 cm to 4.83 cm from the oldest to newest period ($P$<0.0001). Crown root lateral roots increased in length from 3.66 cm to 4.22 cm from 20K to 80K density ($P$=0.0325).

Brace and crown lateral root densities were not affected by Era period. Brace root lateral root density increased 15.8% from 8.42 laterals cm$^{-1}$ to 9.75 laterals cm$^{-1}$ from 20K to 80K ($P$=0.0445), but was not affected by N level. Crown root lateral root density was not affected by any treatment.

Independently, brace and crown root numbers were not significantly affected by Era period. However, the combined brace and crown root number (Fig. 4) decreased from 19.2 in the oldest period to 17.6 in the newest period ($P$=0.0083). Combined root number decreased from 19.4 in HN to 17.4 in
The percentage cortical aerenchyma was not affected by Era period, but increased from 10.5% to 18.8% of cortical area (Fig. 5A) from HN to LN ($P=0.0034$), and from 13.3% to 17% of cortical area from 20K to 80K ($P=0.0026$). The effect of Era period on mean cortical cell size was significant, with size increasing 5% from 0.004 mm$^2$ to 0.0042 mm$^2$ in the newest ($P=0.0232$), but the effects of N level and density were insignificant (Fig. 5B).

Metaxylem vessel number increased from 10.4 in the oldest Era period to 11.9 in the most recent Era period ($P=0.0003$), with no significant effect of N level or density (Fig. 5C). The median area of individual metaxylem vessels decreased 12.2% from 0.00947 mm$^2$ in the oldest material to 0.00831 mm$^2$ in the newest material ($P<0.0001$), with no significant effect of N level or density (Fig. 5D).

**Principal component analysis**

PCA of 15 crown root architectural and anatomical phenes revealed two components that explained 37% of the variation in root phenes (Fig. 6; abbreviations as given in Table 2). The anatomical phenes loaded onto the first component (PC1, 22% of the total variation), and the architectural phenes on the second (PC2, 15% of the total variation). ANOVA including N level, density, and Era as factors indicated no significant effects of these factors on the anatomical component. ANOVA for the architectural component had no significant effect of N level or Era, but the effect of density was significant ($P=0.0041$), with higher density plants having a lower score for PC2.

**Discussion**

**Summary**

Yield of the newest Era maize hybrids surpassed that of the oldest material in every N level and population density combination, and new Era hybrid yield was especially responsive to higher densities in HN relative to the older material, which confirms previous reports (Duvick et al., 2004). High density decreased leaf percentage N, which demonstrates the relationship between population density and N limitation induced by interplant competition. The greater yield of modern Era hybrids relative to older hybrids at greater densities and lower levels of N demonstrates the relatively greater N efficiency of the modern material, as suggested previously in a comparison of two genotypes (Tollenaar and Lee, 2002). To the authors’ knowledge, this is the first study to characterize the root system architecture and root anatomy of US hybrids produced over the past century. The hybrids of the newest Era period were marginally more shallow angled, had fewer nodal roots, had a greater distance to branch, and slightly longer crown root laterals. Anatomically, new Era hybrids had increased cell size and smaller yet more numerous metaxylem vessels. Many of these phene states were previously hypothesized to be optimal for water and nitrate uptake by the SCD ideotype (Lynch, 2013), except nodal root angle needs more careful consideration with regards to the vertical flux of soil sources and its influence on inter- and intraplant competition. Changes in architectural and anatomical root phenotypes in US hybrids over the past 100 years are consistent with the evolution of the maize root system towards integrated phenotypes optimizing N acquisition efficiency in changing agronomic conditions.

**Root system architecture**

The most recent material had brace roots that were 3º more shallow angled and crown roots that were 7º more shallow angles.
angled relative to the oldest material. However, all root angles in this study are steeper than 45° from the horizontal, so may all be classified as relatively steep, especially considering that in the pioneering work of Weaver (1926) many of the angles of outer nodal roots range between 5° and 20° from the horizontal (measured from fig. 84 in that reference). More shallow-angled growth of axial roots has previously been shown to decrease interplant competition for P between...
neighbouring root systems by decreasing the extent of root overlap (Ge et al., 2000), and this logic may be even more important for a more mobile resource such as nitrate where depletion zones caused by root uptake will tend to be larger. Intraplant competition will be decreased when nodal roots are more shallow angled because the proximity of roots of the same plant will also be decreased, as shown with basal root growth angle in common bean (Ge et al., 2000; Rubio et al., 2001). Early in the growing season, the availability of many soil resources, including N and P, is greater in the topsoil (Jobbágy and Jackson, 2001) so shallow-angled roots that explore the topsoil facilitate resource acquisition (Lynch and Brown, 2001). Nitrate leaches in agricultural systems so deep roots may be advantageous later in the season (Di and Cameron, 2002; Thorup-Kristensen, 2006), even though most leaching occurs during fallow periods. Water may also become a deeper resource as the season advances (Asbjørnsen et al., 2008). In common bean, shallow-angled roots that maximized the acquisition of limiting P, a shallow resource, early in the season supported greater plant shoot growth, greater subsequent photosynthesis, greater allocation to the root system, and therefore deeper rooting and greater yield during terminal drought (Ho et al., 2005). Maize with relatively shallow-angled roots that can still grow deeper may co-optimize the acquisition of several soil resources that have differing spatial and temporal availabilities.

The number of nodal roots will determine the overall intensity of soil exploration, and influence the carbon budget of the plant. The newest material had at least 1.6 fewer nodal roots than older lines. Phenes of the outermost brace and crown whorls were measured in this study; however, these are only two of several whorls. The overall decrease in the number of nodal roots could be greater if counted across all nodal root whorls. Recently, Saengwilai et al. (2014b) demonstrated that maize root systems with fewer crown roots maximize N acquisition in LN soil because having fewer nodal roots decreases intraplant root competition, frees carbon and nutrients to be used for other plant tissues, and may allow individual roots to grow longer and explore soil more effectively, which confirmed earlier simulation results (York et al., 2013). Previously, a hybrid released in 1998 was also found to have about one less nodal root per node than a hybrid released in 1959 and to have greater N stress tolerance (McCullough et al., 1994b). Fewer nodal roots may optimize N acquisition while decreasing interplant competition.

The distance from the root tip to lateral emergence may be an important phene for P acquisition in common bean by reducing carbon costs and more efficiently exploring the soil (Miller et al., 2003). In this study, a novel phene termed distance to branching (i.e. distance from the base of the root to the first lateral branch) is reported. For both brace and crown roots, distance to branching almost doubled from the oldest material to the most recent material. The maize root crown contains many nodal roots in close proximity and, by the time the last whorls of crown and brace roots emerge, the soil near the base of the shoot is probably mostly depleted of resources. Delaying the emergence of lateral roots until they are farther from the stem may decrease the carbon costs of lateral root formation while having little or no effect on resource acquisition, which might allow greater carbon allocation to photosynthetic tissue, reproductive tissue, or roots in regions of soil containing greater availability of limiting resources.

Lateral root length has recently been shown to be an important phene in maize for N acquisition (Postma et al., 2014). The most recent material had longer lateral roots on crown roots, but there was no change in lateral root density. Consistent with the change in crown root lateral root length, Postma et al. (2014) demonstrated that fewer but longer lateral roots increase N acquisition when N is limiting, because nitrate is a mobile resource and longer lateral roots expand soil exploration while reducing intraroot competition. Recent field and greenhouse research with several maize genotypes demonstrated a growth advantage for plants with long and few laterals grown in LN soil (Zhan and Lynch, 2015). Thus, the longer crown root laterals observed in the most recent material are consistent with the hypothesis that modern root phenotypes have greater N acquisition efficiency. Possibly, the resources saved by the increased distance to branching in newer material could allow greater expenditures on lateral root length.

**Root anatomy**

Cortical burden reflects the cumulative carbon costs of living cortical tissue (Jaramillo et al., 2013). RCA is formed when cortical cells senesce, leaving behind air spaces which decrease root respiration (Fan et al., 2003). Simulation modelling demonstrated that reduced respiration and remobilization of nutrients after aerenchyma formation lead to increased root length and resource capture (Postma and Lynch, 2011a, b). In the field, maize recombinant inbred lines with greater aerenchyma formation had 800% more yield than lines with less aerenchyma under water stress (Zhu et al., 2010a), and 68% more yield in LN soil (Saengwilai et al., 2014a). In this study, all varieties had more aerenchyma in stressful conditions, such as LN and high density. However, the percentage cortical aerenchyma did not change among Era periods, which could suggest undocumented trade-offs for aerenchyma or a lack of diversity for this phene in the original material. Breeding for more and constitutively expressed aerenchyma may be an important target.

Larger cortical cells are hypothesized to decrease cortical burden by having a lower proportion of cytoplasm volume relative to vacuole volume and less respiratory and nutrient burden (Lynch, 2013). Roots of maize plants with large cortical cell size can respire 59% less than those with small cortical cell size, and can have up to 145% more yield in the field under drought stress (Chimungu et al., 2014a). Cortical cell size increased from the older to the most recent Era periods. Phenes that influence cortical burden may deserve special attention because if they have not been indirectly selected upon, then they may have more potential for yield gains relative to architectural phenes. However, Chimungu et al. (2014a) found a positive relationship between cortical cell size
and leaf cell size, and speculate that genetic determination of plant cell size warrants further study because of possible functional trade-offs in sizes of different cell types. Cortical cell file number (CCFN) is the number of layers of parenchyma cells in the root cortex, and reduced CCFN was associated with greater drought tolerance in maize (Chimungu et al., 2014b); however, the material in this study had no differences in CCFN. Cortical phenes deserve more attention for their possible influences on soil resource acquisition in maize.

All else being equal, smaller diameter metaxylem vessels will decrease water flux (Lewis and Boone, 1995), even if an increase in their number leads to no change in the total area of metaxylem vessels. When water will be limiting during grain filling, conservation of soil water during the early season may increase yield (Lynch et al., 2014), as was observed in a study in which an older maize hybrid used more of the shallow water early in the season than a newer hybrid (Campos et al., 2004). A hybrid with relatively more and smaller diameter xylem vessels than a drought-intolerant hybrid had greater yield under drought stress in the field (de Souza et al., 2013). Cavitation, where the water column is broken in xylem vessels, is problematic, and a hybrid with more xylem vessels had less risk of cavitation (Li et al., 2009), so our observation of more but smaller metaxylem vessels in the newest Era material is consistent with potential drought and cavitation tolerance. Furthermore, fewer nodal roots in newer material may also restrict total water flux and may lead to more productive water use. Root anatomical phenes and their relationships to water and N uptake deserve further attention (Lynch et al., 2014).

Root system evolution

Changes in many of these phenes across Era periods were relatively small. However, even small marginal benefits of many phene states can compound over time to generate measurable performance differences. Duvick et al. (2004) also found small percentage changes for many above-ground phenes in maize, and many of the changes were unintentional, or indirectly selected, as may have happened in the maize root system. In simulations to only 40 d, phenotypes with roots similar to the newest Era plants had the greatest shoot mass because of increased N uptake. Donald (1968) proposed that a cereal ideotype should be a weak competitor. Maize plants with relatively shallow-angled and fewer nodal roots are expected to experience less competition in pure stands than those with extremely steep-angled and many nodal roots, and to avoid root growth redundancy that may negatively affect plant performance (Zhang, 1999). The changes observed in root system architecture and anatomy are consistent with indirect selection for root systems that optimize N acquisition when fields are fertilized with N but high planting densities create competition that leads to individual plants experiencing some level of N stress.

Previous research of root system diversity among individuals of the wild maize ancestor, teosinte, and maize landraces offers further insight into the evolution of the maize root system. In a mesocosm screening of many maize landraces and teosintes, several root architectural phenes were different between landraces and teosintes, while fewer anatomical phenes were different between landraces and teosintes (Burton et al., 2013), which is a similar result to that in this experiment. Anatomical phenes and architectural phenes were also found to form distinct axes in PCA, as in the current study. Architectural phenes may be more readily selected, or anatomical phenes might have stronger constitutive control. Landraces had fewer nodal roots but more seminal roots than teosintes (Burton et al., 2013), which might suggest a general decline in nodal root number as domesticated maize in the USA evolves in more controlled and less limiting environments, as observed in the current study. The reduction in branch points between teosinte and landraces (Burton et al., 2013) may also relate to changes in root system architecture that increase N acquisition efficiency through reduced lateral branching density (Postma et al., 2014). However, in a separate study within maize landraces, increased nodal root number was correlated to increased P acquisition efficiency and also found to be greater in more traditional landraces (Bayuelo-Jimenez et al., 2011), which may demonstrate a trade-off between P acquisition and N acquisition, where more nodal roots are beneficial for growth in low P soil, but fewer nodal roots are beneficial for growth in low N soil. N limitation is more common in US commercial maize fields than P limitation, which may be driving the decrease in nodal root number observed in the newest Era period. In six maize hybrids released between 1973 and 2000 in Northeastern China, root length density decreased in the top 20 cm of soil over time, but remained the same at greater depths (Chen et al., 2013), and these results are consistent with a decrease in nodal root number, especially in the youngest whorls that emerge in shallow soil (Saengwilai et al., 2014). Evidence from the current study and others demonstrates possible evolution of the maize root system mediated by indirect selection towards phene states allowing greater N acquisition efficiency during the original domestication of maize, and its subsequent artificial selection in changing agronomic conditions.

Pioneer hybrids released in each decade from the 1930s to the 1990s have consistently had >40% parentage from the landrace Reid’s Yellow Dent, and the total number of founders was ~60 during this same period (Smith et al., 2004). This diversity is only a fraction of what exists within the genus Zea, and the lack of genetic diversity may relate to a lack of diversity of root phenes. For example, the coefficient of variations (CVs) for nodal root number of the hybrids grown in the 1990s have consistently had >40% parentage from the landrace Reid’s Yellow Dent, and the total number of founders was ~60 during this same period (Smith et al., 2004). This diversity is only a fraction of what exists within the genus Zea, and the lack of genetic diversity may relate to a lack of diversity of root phenes. For example, the coefficient of variations (CVs) for nodal root number of the hybrids grown in the highest density and N levels in this study, landraces, and teosintes in the Burton et al. (2013) study are 0.178, 0.204, and 0.288, respectively. The CV for nodal root number decreases as groups of germplasm go from wild to original domestication to products of the past century’s breeding efforts, which supports that there is, in general, less phenotypic root diversity in elite maize hybrids used in the US than available within the genus Zea as a whole. The introgression of extreme and moderate root phenotype states into elite backgrounds may be a desirable goal for future studies on the utility of specific root phenes.
Conclusion

The trajectories in phenotypic change in this experiment are apparent, and might allow extrapolation to future changes. Breeding has advanced such that more phene states may be introgressed to give rise to plant phenotypes that approach ideotypes for their target environments. Selection for yield and direct selection for specific shoot phenes states have contributed to genetic maize yield gains over the past century (Duvick et al., 2004; Tollenaar and Lee, 2006), but this study demonstrates that changes in both root system architectural and anatomical phenes also occurred over the past 100 years. The changes in the maize root system observed possibly occurred through indirect selection through the effects of root phenes on yield, so direct selection for positive-acting root system phene states could contribute to future yield gains. However, breeding efforts must be based in mechanistic theory that includes how root and shoot phenes will interact and integrate to affect root system resource acquisition and overall plant performance (York et al., 2013). Newer material generated more yield in all agronomic contexts investigated here, and in all contexts, changes in root system phene states across Era periods were consistent with optimizing N acquisition. To the authors’ knowledge, the result of modern material from the Era set generating more yield in LN soil is novel, though McCullough et al. (1994b) found a similar result when comparing a hybrid released in 1959 with a hybrid released in 1988. Possibly, there is less conflict between breeding for high-input and low-input systems than previously recognized due to the intense competition in high-density, high-input systems causing some degree of soil resource stress. Breeding maize with root system architectural and anatomical phene states optimized for N acquisition in limiting conditions has the potential to contribute to more productive and more sustainable systems for both subsistence farmers in developing nations and commercial farmers in developed nations.

Supplementary data

Supplementary data are available at JXB online.
Figure S1. Shoot mass on a per area basis. The RSAJ manual.

Acknowledgements

We thank Dr Joseph Chimungu, Andrew Evensen, Dr Anthony Fernando, Dr Denise Fernando, Elizabeth Nottingham, Hannah Schneider, Robert Snyder, Michael Williams, Jennifer Yang, Ai Zhan, and Xinji Zhang for technical assistance. We thank Dr Carlos Messina and Dr Mark Cooper for their helpful discussions. DuPont Pioneer provided the seed. This research was supported by the National Science Foundation/Basic Research to Enhance Agricultural Development (grant no. 4184-UM-NSF-5380) and Agriculture and Food Research Initiative competitive grant no. 2014-67013-2157 of the USDA National Institute of Food and Agriculture to JPL.

Statement of availability

Plant germplasm and transgenic material will not be made available except at the discretion of the owner and then only in accordance with all applicable governmental regulations. Novel materials described in this publication may be available for non-commercial research purposes upon acceptance and signing of a material transfer agreement. In some cases such materials may contain or be derived from materials obtained from a third party. In such cases, distribution of material will be subject to the requisite permission from any third-party owners, licensors, or controllers of all or parts of the material. Obtaining any permissions will be the sole responsibility of the requestor.

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