



Revista Ciência Agronômica

ISSN: 0045-6888

ccarev@ufc.br

Universidade Federal do Ceará
Brasil

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Revista Ciência Agronômica, vol. 46, núm. 2, abril-junio, 2015, pp. 369-378
Universidade Federal do Ceará
Ceará, Brasil

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Early indirect selection for nitrogen use efficiency in maize¹

Seleção precoce e indireta para eficiência no uso de nitrogênio em milho

Débora Santos Caixeta^{2*}, Roberto Fritsche-Neto³, Ítalo Stefanine Correia Granato⁴, Lucimar Rodrigues de Oliveira⁵ e João Carlos Cardoso Galvão⁶

ABSTRACT - Several studies to evaluate nitrogen use efficiency (NUE) have been carried out using early growth stages. However, there are no scientific reports on the ideal stage for evaluation and on which characteristics have the highest correlation with the NUE at that stage. The aim therefore was to identify the phenological stages and secondary characteristics which maximize accuracy in early indirect selection for NUE in maize. To do this, three endogamic maize strains were evaluated in a completely randomised design with five replications, in a triple factorial scheme (strains x N levels x phenological stage), at two contrasting nitrogen levels: low and high nitrogen. The plants were evaluated at five growth stages: stage nine (V9), with 14 fully-developed leaves (V14), tasseling (VT), flowering (R1) and physiological maturity (R6). The following characteristics were evaluated: efficiency in the usage, absorption, use and translocation of nitrogen; activity of nitrate reductase and glutamine synthetase; length of the lateral and axial roots; specific root area; chlorophyll content; number of leaves; plant height; stem diameter; and the levels of phosphorus and potassium. Considering the estimated direct and indirect gains, it can be concluded that the activities of glutamine synthetase at the V9 and V14 stages permit early indirect selection for nitrogen use efficiency in maize under conditions of low and high N availability respectively.

Key words: *Zea mays*. Path analysis. Abiotic stress. Plant breeding.

RESUMO - Vários trabalhos de avaliação de eficiência no uso de nitrogênio (EUN) já foram feitos utilizando estádios precoces. Contudo, não há relatos científicos sobre o estágio ideal e, neste, quais caracteres que apresentam maior correlação com a EUN. Assim, o objetivo foi identificar os estádios fenológicos e os caracteres secundários que maximizam a acurácia na seleção precoce e indireta para a EUN em milho. Para isso, foram avaliadas três linhagens endogâmicas de milho em delineamento inteiramente casualizado, com cinco repetições, em esquema fatorial triplo (linhagens x níveis de N x estágio fenológico), em duas doses contrastantes de nitrogênio - baixo e alto nitrogênio. As plantas foram avaliadas em cinco estádios fenológicos sendo eles: nove (V9) e 14 folhas completamente desenvolvidas (V14), pendoamento (VT), florescimento (R1) e maturidade fisiológica (R6). Foram avaliados os caracteres: eficiência na utilização, na absorção, no uso e na translocação de nitrogênio, atividade da nitrato redutase e da glutamina sintetase, comprimento de raiz lateral e axial, área de raiz específica, teor de clorofila, número de folhas, altura de planta, diâmetro de colmo, teor de fósforo e potássio. Considerando os ganhos diretos e indiretos estimados, conclui-se que as atividades da glutamina sintetase, nos estádios V9 e V14, permitem a seleção precoce e indireta para eficiência no uso de nitrogênio em milho, nas condições de baixa e alta disponibilidade de N, respectivamente.

Palavras-chave: *Zea mays*. Análise de trilha. Estresse abiótico. Melhoramento vegetal.

DOI: 10.5935/1806-6690.20150016

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¹Recebido para publicação em 19/07/2013; aprovado em 12/12/2014

Parte da Dissertação de Mestrado do primeiro autor apresentada ao Programa de Pós-Graduação em Fitotecnia da Universidade Federal de Viçosa-MG

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INTRODUCTION

Considering only those areas which are suitable for its cultivation, maize production will not be enough to meet world demand for this cereal in the coming decades (FRITSCHENETO; BORÉM, 2011). In addition, approximately 50% of the areas under cultivation are in tropical climates, where conditions of low natural soil fertility and drought predominate (MONNEVEUX *et al.*, 2005). In this context, there are alternatives which can be adopted in order to increase maize production in Brazil, such as the use of off-season crops, the use of cultivars with higher productivity, or even the expansion of farming to lands of the cerrado region. However, some areas of the cerrado impose limitations on agricultural production due to their low soil fertility, especially nitrogen (N). Nevertheless, it is important to note that N is among the main nutrients responsible for increases in maize productivity (RICHARDSON *et al.*, 2009), playing an important part in the metabolism of the plant, especially in protein synthesis (JAKELAITIS; SILVA; FERREIRA, 2005; PAVINATO *et al.*, 2008).

Additionally in recent years, interest in sustainable agriculture has encouraged breeding programs to develop cultivars that show greater efficiency in the use of nutrients, i.e. cultivars displaying satisfactory productivity under conditions of nutritional stress, with consequently lower production costs and environmental impact.

Moll, Kamprath and Jackson (1982) defined nitrogen use efficiency (NUE) as grain or dry-shoot yield per unit of nutrient available to the plant. NUE comprises nitrogen uptake efficiency (NUpE) and nitrogen usage efficiency (NUtE). UpE is the ability of the genotype to absorb the nutrient from the soil, while UtE refers to the ability of the genotype to produce biomass or grains using the absorbed nutrient. Greater values for NUE could therefore be achieved by increasing the UpE and/or UtE (CHEN *et al.*, 2009). According to DoVale *et al.* (2012), regardless of the availability of N, absorption efficiency is the most important component of nitrogen use efficiency. Those researchers also state that regardless of the availability of N, genetic differences as regards NUE are a result of the NUpE. It is therefore important, in breeding programs which aim to increase NUE, for the characteristic used in early selection of superior materials to have a high correlation with the NUpE at the end of the crop cycle. This ensures high productivity and high protein levels in the grain.

The use of methods of early evaluation, or of indirect selection for conditions of abiotic stress, are of great interest as they speed up the selection process, promptly discarding highly susceptible genotypes and enabling resources to be focused on those which are potentially superior (FRITSCHENETO *et al.*, 2010). Because of this, secondary characteristics related to nitrogen use efficiency (NUE) have been used with the

aim of increasing the efficiency of the selection process. In that context, the ideal secondary characteristic is one that is genetically correlated with the NUE under stress, that displays high heritability, has high genetic variability, is easy and quick to evaluate, and is stable during the period of measurement, as well as being observed before flowering, so that there is no recombination of undesirable parents (O'NEILL; SHANAHAN; SCHEPERS, 2006). In this respect, the activities of enzymes related to the metabolism of N can be highlighted (GALLAIS; HIREL, 2004); also the activities of the root system, where root length may be an indicator of the potential for water and nutrient uptake.

Given the above, the objective was to identify the growth stages and secondary characteristics which maximize accuracy in early indirect selection for nitrogen use efficiency in maize.

MATERIAL AND METHODS

Genetic material and conducting the experiment

Three fully endogamous strains of maize were evaluated in a completely randomised design, with five replications, in a triple factorial scheme (strains x levels of N x growth stage), at two contrasting nitrogen dosages: low nitrogen (LN) corresponding to 12.5 mg dm⁻³ urea; and high nitrogen (HN) corresponding to 750 mg dm⁻³ urea. The strains are from the germplasm bank of Programa Milho®, and were previously identified as being efficient, intermediate, and inefficient in the use of nitrogen.

The experiment was set up in a greenhouse at the Federal University of Viçosa, in the state of Minas Gerais, Brazil (MG), from January to May, 2012. Each lot consisted of one plant per pot of 20 litres containing a substrate composed of 50% washed sand and 50% soil from a B horizon of a dystrophic Red-Yellow Latosol, providing negligible amounts of N. Due to this, the values found in soil analysis were disregarded. The seeds were placed into trays to germinate and after five days the best seedlings were transplanted to the pots.

Basic fertilisation was carried out using potassium and phosphorus. Nitrogen fertilisation was by instalments, and followed the nutrient uptake curve of the maize (DUARTE *et al.*, 2003). At each growth stage for which collections were made therefore, enough N was provided to meet plant demand up to that stage only. High availability and stress dosages had been defined in an earlier experiment as 60x the ratio of the two. That experiment was carried out in a completely randomised design, in a 4 x 2 factorial scheme (4 dosages of N x 2 contrasting strains), with two replications; the nitrogen content of the plant being the characteristic evaluated.

Regression analysis was later performed to identify the dosages for high and low nitrogen availability. Low N was considered the dosage at which grain yield in the strains was reduced on average by 50%.

The plants were evaluated (collected) at five growth stages, namely: nine fully developed leaves (V9), 14 fully developed leaves (V14), tasselling (VT), flowering (R1) and physiological maturity (R6).

For each of the five stages being considered, the plants were harvested, the root system was separated from the shoot and the following evaluations performed:

1) Determination of nitrate reductase (NR) activity: by quantifying the nitrite produced per hour, per gram of shoot fresh weight ($\mu\text{moles NO}_2^- \text{ h}^{-1} \text{ g}^{-1} \text{ FW}$). The procedure consists in infiltrating the tissue with a solution containing nitrate, followed by measurement of the nitrite produced by the reaction and which diffuses into the incubation medium. To do this, discs were cut from leaves located on the middle third of the plants. The discs were placed into containers of the incubation medium, which consisted of 0.1 M phosphate buffer pH 7.5, 1% propanol and 0.1 M potassium nitrate. The containers were then incubated in a water bath at 32 °C. During the reaction, the containers were kept in the dark to avoid use of the nitrite formed in the reaction which was catalysed by the NR. After 60 minutes, a 1 ml aliquot part of the reaction medium was removed and placed into a tube containing 2.4 ml of distilled water, 0.3 ml of 1% (w/v) sulfanilamide in 3N HCl, and 0.3 ml 0.02% N-(1-naphthyl) ethylenediamine dichloride. The amount of nitrite (NO_2^-) produced was determined colourimetrically by reading the absorbance of the solutions at 540 nm. The amount of NO_2^- was calculated using a standard curve in the concentration range of 10^{-2} to $10^{-1} \mu\text{mol NO}_2^-$;

2) Determination of glutamine synthetase (GS) activity: by quantifying the glutamyl hydroxamate per hour, per gram of shoot (GSs) and root (GSr) fresh weight ($\mu\text{moles glutamyl hydroxamate (GHD) hour}^{-1} \text{ g}^{-1} \text{ FW}$). The activity can be measured from the synthetase reaction dosing the formation of μ -glutamyl hydroxamate from glutamate and hydroxylamine. The reaction medium used to evaluate enzyme activity by the synthetase reaction was 500 mM glutamate, 60 mM hydroxylamine, 200 mM MgSO_4 , 80 mM ATP and 50 mM imidazole buffer pH 7.4. For the transferase reaction a reaction medium was used which contained 65 mM glutamine, 17 mM hydroxylamine, 33 mM sodium arsenate, 4 mM MnCl_2 , 1.7 mM ADP and 100 mM imidazole buffer pH 6.8. After macerating the leaf discs with the previously frozen roots in the reaction medium, the macerate was placed into an Eppendorf tube and centrifuged at 110,000 rpm for 30 minutes

at 4 °C. After centrifugation, a 1 mL aliquot part of the supernatant was removed and placed into another Eppendorf tube which already contained the reaction medium, and placed into a water bath at 30 °C, the reaction being completed by the addition of the enzyme (0.25 ml). After 20 minutes the reaction was halted by adding 1 ml of a reagent containing 0.67 N HCl, 0.20 N trichloroacetic acid and 0.37 M FeCl_3 . Once the reagent had been added to the supernatant in the test tube, a 2 ml aliquot part was removed, which was again placed into an Eppendorf tube and centrifuged at 100,000 rpm for 5 minutes at 4 °C. The amount of μ -glutamyl hydroxamate formed was then evaluated using a spectrophotometer at a wavelength of 540 nm for the absorbance readings;

3) Determination of shoot dry weight (SDW, g), root dry weight (RDW, g), grain dry weight (GDW, g) and total dry weight (TDW, g). Nitrogen levels and later, nitrogen content, were determined for the shoots (Ns), roots (Nr) and grains (Ng), as well as for total N (Nt). The levels of N were obtained by sulphuric acid digestion of samples of the shoots, roots and grain. Once the data on dry mass and N levels were obtained, and from the amount of nitrogen supplied (Np), efficiency in the absorption, utilisation, use and translocation of nitrogen was estimated according to Moll, Kamprath and Jackson (1982) (Equations 1, 2, 3 and 4):

$$NUE = \frac{TDW}{Nt} \quad (1)$$

$$NUpE = \frac{Nt}{Np} \quad (2)$$

$$NTrE = \frac{Ng}{Nt} \quad (3)$$

$$NUE = NUE \times NUpE = \frac{TDW}{Np} \quad (4)$$

where: NUE is the nitrogen usage efficiency, NUpE the nitrogen uptake efficiency, NUE the nitrogen use efficiency, NTrE the nitrogen translocation efficiency, TDW the total dry weight, Nt the total nitrogen content of the plant, Ng the nitrogen content of the grain, and Np the amount of nitrogen supplied to the plant;

4) Determination of phosphorus levels in the shoots (PLs, dag kg^{-1}), roots (PLr, dag kg^{-1}) and grain (PLg, dag kg^{-1}), and total phosphorus (TP, dag kg^{-1}), as used by DoVale *et al.* (2013);

5) Determination of potassium levels in the shoots (KLs, dag kg^{-1}), roots (KLr, dag kg^{-1}) and grain (KLg, dag kg^{-1}), and total potassium (TK, dag kg^{-1}), as per the method used in determining the P levels;

6) Determination of lateral root length (LRLat, cm), axial root length (LRAXi, cm) and specific root area (SRA, $\text{cm}^2 \text{ g}^{-1}$). Root length and area were obtained with the WinRHIZO 4.1 image analyser, coupled to a

professional Epson XL 10000 Scanner equipped with an additional light unit (TPU). Roots with a diameter less than or equal to 0.5 mm were considered lateral roots, and those over 0.5 mm were considered axial roots (TRASCHSEL *et al.*, 2009). Specific root area was later estimated (SRA, cm² g⁻¹) according to equation 5:

$$SRA = \frac{RA}{RDW} \quad (5)$$

where: RA is the total root area (fine roots + coarse roots) and RDW the root dry weight;

7) Determination of phenological characteristics: number of leaves (NL), plant height (PH, cm), stem diameter (SD cm), chlorophyll levels (SPAD, µg cm⁻²), number of kernel rows per ear (ROW), 1000-grain weight (GW, g) and ear height (EH, cm).

Statistical and genetic analysis

It was found that some characteristics did not have a normal distribution. Data transformation was therefore performed. For the variables NUpE, NUtE and NUE, $\sqrt{(X + 0,5)}$ was used. For LRLat and LRAXi, $\log(x+1)$ was used.

Variance analysis (ANOVA) was later carried out individually (for each level of N within each stage), on the sets from each stage and on the overall set involving all three factors (three strains x two levels of N x five growth stages). In these analyses, the effect of strain (L) was considered to be random, and those of growth stage (S) and availability of N in the soil (N) as fixed. Except for the interaction E x N, all were considered as having a random effect (data not shown).

For those characteristics which were significant by ANOVA, a diagnosis was then made of multicollinearity between the explicative characteristics; their degree in the correlation matrix being established based on condition number. Subsequently, by means of path analysis, breakdowns were made of the phenotypic correlations for low N availability, for the direct and indirect effects of the characteristics LRLat, LRAXi, SRA, NR, GSs, GSr, SPAD, NL, PH, SD, TP, TK, NTrE (explicative variables) on the characteristic NUE and its components NUpE and NUtE (dependent variables), according to their significance at each stage under evaluation (tables not shown). Finally, gains were estimated for the dependent variables at the two levels of N, with direct (GSd) and indirect selection (GSi) of the explicative variables.

The analyses were performed using the Genes Software for Genetics and Experimental Statistics (CRUZ, 2013) and the Statistical Analysis System (SAS) version 9.1 (SAS INSTITUTE, 2003).

RESULTS AND DISCUSSION

Path analysis

By means of path analysis, it was sought to quantify the effect of the secondary characteristics during the growth stages being analysed, on the NUpE at the R6 stage.

At stage V9, under low N availability, a high direct effect was seen from the GSs on the NUpE at R6 (2.55) in relation to the residual effect (0.01) (Table 1). This indicates that plants with high GSs activity early in the cycle will present high values for NUpE at the end of the cycle, with a consequently high NUE. High GSs activity makes the process of N assimilation more efficient. Signals are sent to the plant that the absorption process should continue as there is no accumulation of NH₄⁺. This could generate negative feedback, reducing the absorption rate, besides being a toxic substrate for the plants at high concentrations. From the high coefficient of determination (0.99), it was also seen that the characteristics GSs, LRAXi and TP explain almost the total variation in NUpE.

In addition to GSs, the characteristic LRAXi was also potentially efficient in the early and indirect selection for NUpE at R6 due to its high direct effect (1.63). However, the use of the two characteristics GSs and LRAXi simultaneously in order to increase the accuracy of the selection process, is not possible, as they have negative indirect effects on each other (-1.39 and -2.17). Increasing the GSs is therefore the most appropriate method because it is not destructive, allowing the plants that were evaluated to intercross, a situation which is not possible when evaluating plants for LRAXi.

At stage V14 under low N availability, the characteristics TP and TK explain all the phenotypic variation for the NUpE at R6 (R² = 1.00) (Table 2). For indirect selection at the beginning of the cycle, it is seen that genotypes with high levels of P and low levels of K are correlated with a high NUpE at the end of the cycle (0.61). This high negative correlation of K with NUpE lies in the fact that excess potassium restricts N uptake. However, it is known that K is related to the activity of the enzyme NR, due to being associated with the regulation and uptake of nitrate by the plant roots. According to Silva *et al.* (2011), when nitrate is reduced in the leaves, malate is formed in response to the increased internal pH of the tissue. According to the same authors, part of the malate is transferred to the phloem, accompanied by potassium, to be translocated to the roots, where it undergoes decarboxylation. The carbonic acid produced is released into the environment in exchange for the absorbed nitrate, in that way controlling the absorption of the nitrate. The recirculated potassium serves as a counterion for the transport of nitrate in the xylem to the shoots. It is therefore of great importance to maintain the K content of the substrate at optimal levels, so that there is no negative interference with the absorption of N.

The relation of P to NUpE has been demonstrated in several studies. Among them, Alves *et al.* (1996) observed a substantial reduction in levels of total N in maize plants exposed to increasing periods of P omission in a nutrient solution. This is due to the importance of P as a constituent of adenosine triphosphate (ATP), generating the necessary energy for the activation of GS, since the assimilation of this nutrient is a highly demanding process in terms of energy, requiring the transfer of two electrons per NO_3^- converted into NO_2^- , six electrons per NO_2^- converted into NH_4^+ and two electrons

and one ATP per molecule of NH_4^+ converted into glutamate (BLOOM; SUKRAPANNA; WARNER, 1992).

At the tasselling stage (VT) under low N availability, the presence of severe multicollinearity was diagnosed between the explicative characteristics related to the condition of low N availability in the phenotypic correlation matrix for the path analysis. Since all the characteristics are potentially important and cannot be eliminated, path analysis for this level of N was performed under multicollinearity (CARVALHO; CRUZ, 1996).

Table 1 - Estimates of the direct and indirect effects between the main dependent variable, NUpE at stage R6, and the independent explicative variables, GSs, LRAXi and TP, in three strains of maize evaluated under low nitrogen availability (LN), at stage V9, Viçosa, MG, Brazil

Characteristic	Effects of Association	Estimate
GSs ²	Direct effect on NUpE ¹	2.55
	Indirect effect through LRAXi	-1.39
	Indirect effect through TP	-1.33
Total		-0.17
LRAXi ³	Direct effect on NUpE	1.63
	Indirect effect through GSs	-2.17
	Indirect effect through TP	1.20
Total		0.66
TP ⁴	Direct effect on NUpE	-1.34
	Indirect effect through GSs	2.54
	Indirect effect through LRAXi	-1.47
Total		-0.26
Coefficient of Determination		0.99
Effect of residual variable		0.01

¹ Nitrogen uptake efficiency. ² Glutamine synthetase activity in the shoots ($\mu\text{moles GHD h}^{-1} \text{g}^{-1} \text{FW}$). ³ Axial root length (cm). ⁴ Level of phosphorus (dag kg^{-1})

Table 2 - Estimates of the direct and indirect effects between the main dependent variable, NUpE at stage R6, and the independent explicative variables, TP and TK in three strains of maize evaluated under low nitrogen availability (LN), at stage V14, Viçosa, MG, Brazil

Characteristic	Effects of Association	Estimate
TP ²	Direct effect on NUpE ¹	0.76
	Indirect effect through TK	-0.15
Total		0.61
TK ³	Direct effect on NUpE	-0.80
	Indirect effect through TP	0.14
Total		-0.66
Coefficient of Determination		1.00
Effect of residual variable		0.00

¹ Nitrogen uptake efficiency. ² Level of phosphorus (dag kg^{-1}). ³ Level of potassium (dag kg^{-1})

The value for the constant k chosen for stabilisation of the estimates was relatively small (0.05), producing little bias in the regression analysis. This can be seen by the high coefficient of determination of the path analysis ($R^2 = 0.98$), showing that most of the variation in the principle characteristic was determined by the explicative characteristics (Table 3). It was seen that correlation was negative (-0.99) between the NUtE and the NUpE, possibly because the plant prioritises one component over the other during its development. This is because maize plants in the early stages of development probably absorb more nutrients than they use. The NUpE therefore becomes the most important component in these early stages of the cycle. However, in the course of development, the use component sees an increase in importance at the expense of uptake. The negative direct effects of LRAXi (-0.22) and LRLat (-0.23) are contrary to the expected increase in N uptake from the increase in root length. This may be due to the plant expending an excess of energy on root growth, decreasing availability for absorption.

At this stage (VT), it is also known that GSr has both a moderate direct effect (0.14) on NUpE and a high correlation (0.64), and can also be used as a characteristic in preparing selection indices for plants with high NUpE. This demonstrates that assimilation of N occurs not only in the leaves, but also in the roots. However, for this to occur, it is necessary to import carbohydrates produced in the leaves, so that after being metabolised they can generate energy and provide carbon skeletons for the process of nitrogen

incorporation into amino acids (OAKS, 1994), the process expending more energy compared to assimilation in the leaves. This energy advantage is the basis of the suggestion for selecting strains that maximise the reduction of NO_3^- in the leaves, with a view to reducing energy requirements and increasing productivity. One proviso that must be made, is that from this stage on, there is no longer a way to maintain only superior plants for hybridisation, thereby reducing selection gain. In addition, the characteristics that explained the NUpE are destructive, hindering the process of rapid evaluation, and making it impossible for the materials evaluated to be used as units for recombination. There will consequently be a reduction in any potential gains to be made from the selection.

At the R1 stage under low N availability, the characteristics considered in the path analysis also explain almost all of the total variation in NUpE ($R^2 = 0.98$) (Table 4). For this case there was also severe multicollinearity in the phenotypic correlation matrix, the path analysis being performed under multicollinearity with the value of k equal to that in the previous analysis (0.05). It was seen that the direct effect of GSs on the NUpE decreased when compared to the V9 stage (0.33). However, there was an increase in correlation (0.87), reconfirming the significance of the interaction, strain x evaluation stage. Root length had a direct effect on the NUpE similar to that seen at the VT stage.

Table 3 - Estimates of the direct and indirect effects between the main dependent variable, NUpE at stage R6, and the independent explicative variables, NUtE, NUE, LRLat, LRAXi, NL and GSr in three strains of maize evaluated under low nitrogen availability (LN), at stage VT, Viçosa, MG, Brazil

Characteristic	Effects of Association	Estimate	VIF*
NUtE ²	Direct effect on NUpE ¹	-0.23	16.80
	Indirect effect through NUE	-0.23	15.22
	Indirect effect through LRLat	-0.23	14.87
	Indirect effect through LRAXi	-0.22	14.81
	Indirect effect through NL	-8E-4	0.07
	Indirect effect through GSr	-0.08	3.94
Total		-0.99	
NUE ³	Direct effect on NUpE	-0.23	16.82
	Indirect effect through NUtE	-0.23	15.20
	Indirect effect through LRLat	-0.23	14.76
	Indirect effect through LRAXi	-0.22	14.69
	Indirect effect through NL	-5E-4	0.03
	Indirect effect through GSr	-0.08	4.28
Total		-0.99	

Table 3 Continuation

LRLat ⁴	Direct effect on NUpE	-0.23	16.58
	Indirect effect through NUtE	-0.23	15.06
	Indirect effect through NUE	-0.23	14.97
	Indirect effect through LRAxi	-0.22	15.01
	Indirect effect through NL	-2E-3	0.29
	Indirect effect through GSr	-0.07	2.90
Total		-0.98	
LRAxi ⁵	Direct effect on NUpE	-0.22	16.57
	Indirect effect through NUtE	-0.23	15.02
	Indirect effect through NUE	-0.22	14.91
	Indirect effect through LRLat	-0.23	15.02
	Indirect effect through NL	-2E-3	0.34
	Indirect effect through GSr	-0.07	2.77
Total		-0.98	
NL ⁶	Direct effect on NUpE	-0.01	8.96
	Indirect effect through NUtE	-0.02	0.12
	Indirect effect through NUE	-0.01	0.05
	Indirect effect through LRLat	-0.04	0.55
	Indirect effect through LRAxi	-0.05	0.62
	Indirect effect through GSr	0.10	6.24
Total		-0.03	
GSr ⁷	Direct effect on NUpE	0.14	12.32
	Indirect effect through NUtE	0.14	5.37
	Indirect effect through NUE	0.14	5.84
	Indirect effect through LRLat	0.12	3.91
	Indirect effect through LRAxi	0.11	3.73
	Indirect effect through NL	-0.01	4.53
Total		0.64	
Coefficient of Determination		0.98	
Value of k		0.05	
Effect of residual variable		0.11	

* Variance inflation factor. ¹ Nitrogen uptake efficiency. ² Nitrogen usage efficiency ³ Nitrogen use efficiency. ⁴ Lateral root length (cm). ⁵ Axial root length (cm). ⁶ Number of leaves. ⁷ Glutamine synthetase activity in the roots ($\mu\text{moles GHD h}^{-1} \text{g}^{-1} \text{FW}$)

Table 4 - Estimates of the direct and indirect effects between the main dependent variable, NUpE at stage R6, and the independent explicative variables, GSs, LRLat, LRAxi and NL in three strains of maize evaluated under low nitrogen availability (LN), at stage R1, Viçosa, MG, Brazil

Characteristic	Effects of Association	Estimate	VIF*
GSs ²	Direct effect on NUpE ¹	0.33	12.01
	Indirect effect through LRLat	0.30	8.22
	Indirect effect through LRAxi	0.20	4.62
	Indirect effect through NL	5E-3	1.56
Total		0.87	

Table 4 Continuation

LRLat ³	Direct effect on NUpE	-0.37	13.53
	Indirect effect through GSs	-0.27	7.29
	Indirect effect through CRAxi	-0.34	10.63
	Indirect effect through NF	-7E-4	0.03
Total		-0.99	
LRAxi ⁴	Direct effect on NUpE	-0.35	12.65
	Indirect effect through GSs	-0.21	4.39
	Indirect effect through LRLat	-0.36	11.38
	Indirect effect through NL	3E-3	0.68
Total		-0.93	
NL ⁵	Direct effect on NUpE	-0.01	6.58
	Indirect effect through GSs	-0.17	2.86
	Indirect effect through LRLat	0.03	0.07
	Indirect effect through LRAxi	0.12	1.31
Total		-0.03	
Coefficient of Determination		0.98	
Value of k		0.05	
Effect of residual variable		0.14	

* Variance inflation factor. ¹ Nitrogen uptake efficiency. ² Glutamine synthetase activity in the roots ($\mu\text{moles GHD h}^{-1} \text{g}^{-1} \text{FW}$). ³ Lateral root length (cm). ⁴ Axial root length (cm). ⁵ Number of leaves

Gains from selection

Considering the direct and indirect gains estimated for NUpE at stage R6 under LN, it can be seen that the greatest gain would be obtained by selecting for GSs at stage R1 - a fairly advanced stage in the plant cycle (Table 5). However, it is of great interest that the evaluation be made early, as this speeds up the selection process, with the least efficient genotypes being discarded immediately and consequently only superior individuals being interbred. (FRITSCHENETO *et al.*, 2010). It therefore follows that GSs at V9, with an estimated gain of 83.55%, is the most promising characteristic for indirect, early selection in this environment. In addition to its high heritability (82.9%, data not shown), as the method is non-destructive, it is possible to evaluate the recombination units themselves, maximizing control over the parents and the gains when selecting for NUpE at the end of the cycle.

Under HN, GSr at stage V14 was the characteristic which made possible a greater indirect gain of the NUpE in a stress environment at the end of the cycle (206.86%) (Table 6). In addition, selection at stage V14 makes it possible to keep only superior individuals for the recombination process. In an HN

environment, there is also the advantage of the plants fully expressing their genetic potential, with the possibility of evaluating other variables of interest to commercial farming, such as lodging and resistance to disease.

The enzyme NR was not so important in the selection of superior genotypes as the NUE. Similar results were observed by Oliveira *et al.* (2013), where the activity of the enzyme did not differentiate between varieties which were efficient or inefficient in the use of N. In the present work, NR activity was also not influenced by N levels. This shows that high NR activity does not necessarily result in high NUE, seeing that NR is only the first enzyme in a long process of assimilation. On the other hand, GSs is the characteristic with the greatest effect on NUpE, and is consequently of interest in the evaluation of genotypes for NUE. Some authors (FONTAINE *et al.*, 2009; MARTIN *et al.*, 2006) have confirmed that quantitative trait loci (QTL) identified for grain yield and its components, coincide with QTL identified for GS activity. Furthermore, it has been demonstrated in maize and rice mutants that GS is necessary for grain filling in the presence and absence of N (HIREL *et al.*, 2007). Thus, selecting for GSs activity will result in plants which are more efficient in the use of nitrogen and hence have higher yields.

Table 5 - Estimates of direct (GSd) and indirect (GSI) gains (%) with selection for the characteristics: GSs, LRAXi and TP at stage V9; TP and TK at stage V14; NUtE, NUE, LRLat, LRAXi and GSs at stage VT; and GSs, LRLat and LRAXi at stage R1, on NUpE at stage R6, in three strains of maize evaluated under low nitrogen availability (LN), in Viçosa, MG, Brazil

Selection	Selection Stage	Selection Characteristic	Response Characteristic	Response Stage	GS%
Indirect	V9 ⁴	GSs ¹	NUpE	R6	83.55
		LRAXi ²			40.19
		TP ³			27.25
Indirect	V14 ⁶	TP	NUpE	R6	43.92
		TK ⁵			34.60
Indirect	VT ¹¹	NUtE ⁷	NUpE	R6	53.49
		NUE ⁸			125.42
		LRLat ⁹			-0.21
		LRAXi			-3.26
		GSr ¹⁰			158.60
Indirect	R1 ¹²	GSs	NUpE	R6	179.44
		LRLat			2.63
		LRAXi			-1.04
Direct	R6 ¹⁴	NUpE ¹³	NUpE	R6	10.91

¹Glutamine synthetase activity in the shoots ($\mu\text{moles GHD h}^{-1} \text{g}^{-1} \text{FW}$). ² Axial root length (cm). ³ Phosphorus content (dag kg^{-1}). ⁴ Nine fully developed leaves. ⁵ Potassium content (dag kg^{-1}). ⁶ Fourteen fully developed leaves. ⁷ Nitrogen usage efficiency. ⁸ Nitrogen use efficiency. ⁹ Lateral root length (cm). ¹⁰ Glutamine synthetase activity in the roots ($\mu\text{moles GHD h}^{-1} \text{g}^{-1} \text{FW}$). ¹¹ Tasselling. ¹² Flowering. ¹³ Nitrogen uptake efficiency. ¹⁴ Physiological maturity

Table 6 - Estimates of direct (GSd) and indirect (GSI) gains (%) with selection for the characteristics: NUpE, NUE, CRAXi, TP and TK at stage V9; NUpE, NUE, GSs, LRLat, LRAXi and TP at stage V14; NUpE, LRLat, LRAXi, TP and TK at stage VT; and TP at stage R6, on NUpE at stage R6, in three strains of maize evaluated under high nitrogen availability (HN), in Viçosa, MG, Brazil

Selection	Selection Stage	Selection Characteristic	Response Characteristic	Response Stage	GS%
Indirect	V9 ⁶	NUpE ¹	NUpE	R6	75.71
		NUE ²			14.57
		LRAXi ³			9.69
		TP ⁴			28.76
		TK ⁵			12.50
Indirect	V14 ⁹	NUpE	NUpE	R6	33.21
		NUE			13.05
		GSs ⁷			206.86
		LRLat ⁸			6.28
		LRAXi			12.01
Indirect	VT ¹⁰	TP	NUpE	R6	39.39
		NUpE			151.43
		LRLat			37.96
		LRAXi			30.80
		TP			41.77
Indirect	R6 ¹¹	TK	NUpE	R6	34.81
		TP			27.01
Direct	R6 ¹¹	NUpE	NUpE	R6	0.23

¹ Nitrogen uptake efficiency. ² Nitrogen use efficiency. ³ Axial root length (cm). ⁴ Phosphorus content (dag kg^{-1}). ⁵ Potassium content (dag kg^{-1}). ⁶ Nine fully developed leaves. ⁷ Glutamine synthetase activity in the shoots ($\mu\text{moles GHD h}^{-1} \text{g}^{-1} \text{FW}$). ⁸ Lateral root length (cm). ⁹ Fourteen fully developed leaves. ¹⁰ Tasselling. ¹¹ Physiological maturity

CONCLUSION

The activities of glutamine synthetase at stages V9 and V14 allow early indirect selection for nitrogen use efficiency in maize under conditions of low and high N availability respectively.

ACKNOWLEDGEMENT

The authors wish to thank FAPEMIG for a scholarship grant and for funding the research project (CAG - APQ-00559-12).

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