

Performance of low phosphorus tolerant rice genotypes under drought stress

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ABSTRACT

The identification of genotypes more tolerant to water deficit and more efficient in the use of nutrients little available in the soil is an important low-cost strategy to promote sustainable agriculture in marginalized regions. In this sense, a study was carried out in the SITIS Phenotyping Platform of the Embrapa Arroz e Feijão, with the objective of evaluating the drought tolerance of upland rice genotypes tolerant to phosphorus (P) deficiency. The experimental design was performed in a randomized complete block with split-split plots and two replications. In the plots were established two water regimes (with and without water deficit), in the subplots two soil phosphorus contents (25 and 200 mg dm⁻³) and in sub-subplots 48 upland rice genotypes. The water deficit had more impact on grain yield than the phosphorus supply. The genotypes AB 062037, AB 062041, AB 062138, Arroz Mato Grosso, BRA 02601, BRA 052045, CNA 4098, CNA 6187, Guapa, Guaporé and Rio Paranaíba were classified in the most productive group under both water regimes. The most productive genotypes under water deficit showed higher root density in the deeper soil layers. The most productive genotypes in the two water regimes were also those that showed the highest transpiration.

Keywords: Oryza sativa L.; abiotic stress; root system; evapotranspiration.

RESUMO

Desempenho de genótipos de arroz tolerantes a baixo nível de fósforo sob deficiência hídrica

A identificação de genótipos mais tolerantes à deficiência hídrica e mais eficientes no uso de nutrientes pouco disponíveis no solo é uma importante estratégia de baixo custo para promover a agricultura sustentável em regiões marginalizadas. Neste sentido, foi conduzido um estudo na Plataforma SITIS de Fenotipagem da Embrapa Arroz e Feijão, com o objetivo de avaliar a tolerância à deficiência hídrica de genótipos de arroz de terras altas tolerantes à deficiência de fósforo. O delineamento experimental foi blocos ao acaso, com parcelas subsubdivididas, com duas repetições. Nas parcelas foram estabelecidos dois regimes hídricos (com e sem deficiência hídrica), nas subparcelas dois teores de fósforo no solo (25 e 200 mg dm⁻³) e nas subsubparcelas 48 genótipos de arroz de terras altas. A deficiência hídrica teve mais impacto na produtividade de grãos do que o suprimento de fósforo. Os genótipos AB 062037, AB 062041, AB 062138, Arroz Mato Grosso, BRA 02601, BRA 052045, CNA 4098, CNA 6187, Guapa, Guaporé e Rio Paranaíba foram classificados no grupo mais produtivo sob ambos regimes hídricos. Os genótipos mais produtivos sob deficiência hídrica apresentaram maior densidade do sistema radicular nas camadas mais profundas do solo. Os genótipos mais produtivos nos dois regimes hídricos foram também aqueles que apresentaram maior transpiração.

Palavras-chave: Oryza sativa L.; estresse abiótico; sistema radicular; evapotranspiração.

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INTRODUCTION

Upland rice is grown in almost all regions of Brazil and in a wide range of climatic conditions and production systems, covering from large mechanized areas to production systems for subsistence. It is characterized as a sensitive crop to drought stress. According to Guimarães *et al* (2016), this situation should worsen with global warming, because the water deficit should be strengthened in areas where it already occurs and emerge in other regions that are not currently subject to this stress.

In the Brazilian savannahs region, where the upland rice is mostly grown, the soils showed low available water capacity, low phosphorus and high aluminum content, which are limiting factors for yield. The availability of phosphorus (P) applied as fertilizer is generally limited, due to the abundance of iron and aluminum oxides in these soils, which makes the phosphorus nutrition a limiting factor to achieving economically satisfactory farm incomes (Costa *et al.*, 2006; Crusciol *et al.*, 2006). In addition, phosphates have achieved high international prices, which has burdened the cost of upland rice production, since we are dependent on its importation.

Considering these aspects, the identification of genotypes more tolerant to water deficit and more efficient in the use of nutrients little available in the soil is an important low-cost strategy to promote sustainable agriculture in marginalized regions (Otani & Ae, 1996). Wissuwa & Ae (2001) confirmed this strategy by transferring the major quantitative trait locus (*Phosphorus uptake1* - Pup1) of high capacity of P uptake of the cultivar 'Kasalath', tolerant to P deficiency, for modern rice cultivar 'Nipponbare', with high harvest index, which tripled its grain yield in conditions of low P availability.

Chin *et al.* (2010) showed that Pup1 is present in more than 50% of rice accessions adapted to stress-prone environments, whereas it was detected in only about 10% of the analyzed irrigated/lowland varieties. Furthermore, the Pup1 locus was detected in more than 80% of the analyzed drought tolerant rice breeding lines.

A deep root system could improve the adaptation of rice during drought through greater capacity for water extraction, thus maintaining high plant leaf water status (Kamoshita *et al.*, 2004). In addition, a well-developed root system, able to exploit large soil volume, is recognized as one of the most important plant adaptation mechanisms to ensure adequate nutrient uptake (Yang *et al.*, 2004).

In this sense, it was tested the hypothesis that genotypes more efficient in the use of phosphorus are also tolerant to the water deficit and that those with higher root density in depth are more adapted to water deficit periods and low soil P availability. Therefore, this study aimed to identify, among rice genotypes tolerant to P deficiency, those with tolerance to drought stress, as a low-cost strategy to produce rice in regions of Brazilian savannahs subject to these stresses.

MATERIAL AND METHODS

The study was carried out in a greenhouse, in soil columns, at Embrapa Arroz e Feijão' SITIS Phenotyping Platform, in Santo Antônio de Goiás, GO. The soil used was an Oxisol, whose chemical analysis showed the following results: pH (H₂O) = 5.1; Ca = 5.0 mmolc dm⁻³; Mg = 4 mmol_c dm⁻³; Al = 2 mmol_c dm⁻³; P = 0.4 mg dm⁻³; K = 39 mg dm⁻³; Cu = 1.7 mg dm⁻³, Zn = 1.1 mg dm⁻³, Fe = 29 mg dm⁻³, Mn = 19 mg dm⁻³ and organic matter = 35.2 g kg⁻¹.

The SITIS Phenotyping Platform is a real-time automated control system for monitoring plant physiological parameters, soil moisture, greenhouse climate and irrigation of the soil columns. It is composed of 384 soil columns, packed in PVC pipes with a 0.25 m diameter and 1.00 m height, formed of five rings 0.20 m high connected by duct tape, and placed on digital scales with an irrigation point for each column. The amount of water used by the plants can be monitored in each column by the difference in weight.

The experimental design was performed in a randomized complete block with split-split plots and two replicates. In the plots were established two water regimes, in the subplots two soil phosphorus contents and in sub-subplots 48 upland rice genotypes from different origins. Among these, 47 genotypes were tolerant to P-deficient soils by preliminary field trial, and one was not tolerant (Farroupilha), used as a negative check.

In one of the water regimes, genotypes were kept in adequate soil moisture conditions throughout the crop cycle, -0.025 MPa at 0.1 m depth (Pinheiro *et al.*, 2006). In the other, they were kept under these conditions until the beginning of panicle emission (R3 stage), when they were subjected to water deficit, which was maintained until the end of the crop cycle, with daily replacement of 50% of the water lost by evapotranspiration, which was monitored by weighing. The soil P contents were 25 and 200 mg dm⁻³, stablished in the soil layer of 0-0.2 m depth with necessary supplementation with triple superphosphate.

Three seeds were sown per column and were thinned seven days after emergence, leaving one plant per column. In addition to P fertilization, 1.33 g of KCl and 1.0 g of ammonium sulfate per soil column were applied at sowing and 2 g of ammonium sulfate per column at topdressing, 45 days after emergence.

The root system was evaluated at the time of the grain harvest in 0.2 m soil layers from the surface to 1 m deep. The separation of the roots from the soil samples was performed by the method of repeated suspension/ decantation. After separation, the roots were recovered from the supernatant in 0.25 mm sieves with a tweezer, and taken to a forced air-circulating oven at 80°C until a constant weight was obtained, to determine its dry matter weight. The root density was obtained by dividing its dry biomass, in mg, by the sample volume, in cm³.

Grain yield and dry biomass of leaves and stems per column, in g, and evapotranspiration after the panicle emission stage, in L per column, obtained by the daily weighing of soil columns, were also determined. Data were submitted to Shapiro-Wilk test to verify the normality of variances. Since the data were parametric, the analysis of variance was used and the means were compared by Scott-Knott's test. Correlations among all variables were also performed.

RESULTS AND DISCUSSION

Water regimes significantly affected all agronomic traits of the genotypes, except the root density in the 0-0.2 m layer (Table 1). The average grain yields obtained with and without water deficit were 33.4 and 91.0 g per column, the leaf biomasses were 23.8 and 29.3 g per column and the stem biomasses were 46.9 and 61.3 g per column, respectively (Tables 2 and 3). Therefore, the water deficit induced a significant reduction of 63.3%, 18.8% and 23.5% in grain yield and biomass of leaves and stems, respectively.

Water deficit also induced reductions in root density, which were of 13.1%, 13.3%, 10.5%, 23.2% and 29.1% in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth, respectively (Tables 2 and 3).

The single effect of the soil P content was not significant for any of the agronomic traits of the genotypes

probably due to the genotypes were tolerant to P-deficient soils, except one (Farroupilha), that was the negative check. Under water deficit, this genotype showed the lowest grain yield in absolute value (Table 2) and, under well-watered conditions, was in the least productivity group (Table 3).

The water regime x soil P content interaction was significant for grain yield (Table 1). In the presence of water deficit, the grain yield at the lowest P content was 36.1 g per column, 17.6% significantly higher than that obtained with the highest P content, which was 30.7 g per column. In the absence of water deficit, instead, a 9.8% higher yield, 95.3 g per column, was obtained with the highest P content, compared with 86.8 g per column obtained at the lowest P content.

According to Dingkuhn et al. (2006), the main effect of P deficiency appeared to be a reduction in demand for assimilates in the shoot while photosynthetic radiation use efficiency remained nearly constant; resulting in spillover of excess assimilates into reserve compartments and root growth. Increased root growth relative to shoot was associated with increased sucrose concentration in roots, and thus possibly resulted from assimilates liberated by shoot growth inhibition (Luquet et al., 2005). According to Wissuwa et al. (2005), roots are generally regarded as stronger sinks than leaves under P deficiency since root:shoot ratios typically increase. Additionally, in the comparison of two genotypes with different tolerance to P deficiency, these authors found that the more tolerant one preferentially distributed P to roots where the additional P stimulated root growth and, ultimately, P uptake.

In our study, under water deficit, the average root density in the 0.4-1.0 m deeper layers was 0.50 mg cm⁻³

Source of variation	DF	Mean square								
		GY	LeafBio	StemBio -	Root density					
					0-0.2 m	0.2-0.4 m	0.4-0.6 m	0.6-0.8 m	0.8-1.0 m	
Replication	1	975.1 ^{ns}	5.8 *	230.2 ns	0.32 ns	0.12 ns	0.17 ^{ns}	0.10 ns	0.06 ns	
Water regime (WR)	1	318,718.1 **	2,896.8 **	19,966.4 *	1.03 ns	0.50 **	0.28 **	1.81 **	4.26 *	
Error A	1	62.5	0.005	39.5	0.05	0.01	0.01	0.09	0.45	
P content (P)	1	243.8 ns	363.0 ns	3,786.1 ^{ns}	0.07 ns	0.25 ns	0.02 ns	0.03 ns	0.10 ns	
WR x P	1	4,634.3 **	276.0 ns	1,179.6 ^{ns}	0.01 ns	0.01 ns	0.01 ns	0.17 ^{ns}	0.53 ns	
Error B	2	160.7	32.2	583.1	1.23	0.12	0.03	0.03	0.05	
Genotype (G)	47	1,031.7 **	463.3 **	1,331.3 **	0.62 **	0.17 **	0.11 **	0.12 **	0.19 **	
WR x G	47	435.8 **	22.2 ns	117.2 *	0.08 ns	0.05 **	0.05 *	0.04 **	0.12 **	
P x G	47	200.1 ns	23.0 ns	84.7 ns	0.05 ns	0.03 ns	0.04 ns	0.03 ns	0.07 ns	
WR x P x G	47	214.9 ns	21.1 ns	80.1 ns	0.05 ns	0.03 ns	0.02 ns	0.03 ns	0.05 ns	
Error C	188	186.8	21.1	81.8	0.08	0.03	0.03	0.02	0.05	
CV (%)		22.0	17.3	16.7	37.6	33.6	36.8	29.9	38.2	

Table 1: Summary of variance analysis for grain yield (GY, g per column), leaf biomass (LeafBio, g per column), stem biomass (StemBio, g per column) and root density (mg cm⁻³) in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth

^{ns}, * and **: not significant, significant at 0.05 and 0.01 levels by F-test, respectively.

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Constune	GY	LeafBio	StemBio		Root	density (mg	cm ⁻³)	
Genotype	(g per colum	n)	0-0.2 m	0.2-0.4 m	0.4-0.6 m	0.6-0.8 m	0.8-1.0 m
CNA 4140	51.6 a	35.5 a	59.6 c	1.06 b	0.64 a	0.65 a	0.68 a	0.95 a
CNA 6187	50.3 a	26.2 c	54.8 c	0.88 b	0.44 b	0.56 a	0.45 b	0.51 b
Arroz Mato Grosso	50.0 a	31.7 b	58.0 c	0.76 b	0.61 a	0.78 a	0.64 a	0.66 a
BRA 02601	45.9 a	36.1 a	66.9 c	0.85 b	0.62 a	0.63 a	0.79 a	0.85 a
BRS Esmeralda	44.8 a	20.3 d	36.9 e	0.35 c	0.43 b	0.41 b	0.46 b	0.41 b
BRA 052023	44.8 a	23.6 c	44.8 d	0.60 c	0.52 b	0.50 a	0.46 b	0.42 b
Rio Paranaíba	44.4 a	39.7 a	62.0 c	0.83 b	0.40 b	0.51 a	0.64 a	0.88 a
TOX 503-4-115-B-B	43.0 a	25.3 c	46.9 d	1.04 b	0.61 a	0.39 b	0.56 a	0.64 a
BRS Bonanca	42.8 a	21.9 c	40.1 d	0.96 b	0.50 b	0.42 b	0.46 b	0.64 a
Guaporé	42.4 a	36.3 a	47.9 d	0.89 b	0.84 a	0.55 a	0.48 b	0.79 a
CNA 4137	41.1 a	13.1 d	33.7 e	0.38 c	0.20 h	0.28 b	0.32 b	0.29 b
CNA 4098	40.8 a	30.0 b	47.5 d	1.08 b	0.74 a	0.44 b	0.54 a	0.88 a
Guana	40.8 a	36.0 0 36.7 a	64.4 c	0.91 b	0.53 h	0.61 a	0.54 a	0.00 a
BRA 052045	40.3 a	30.7 a	68.1 c	1 45 a	0.65 a	0.01 a	0.02 u	0.73 a
AB 062041	40.1 a	30.2 h	53.9 c	1.43 a 1.03 b	0.68 a	0.50 a	0.420	0.75 a
AB 062037	30 8 a	30.1 b	64.1 c	0.84 b	0.00 a	0.37 h	0.07 a	0.77 a
IDEM 247	39.0 a	15.8 d	40.3 d	0.840	0.47 b	0.57 0	0.48 U	0.30 D
IKEWI 247	37.0 a	13.6 d	40.3 u	0.43 C	0.45 b	0.50 a	0.37 b	0.47 U
DDSMC 255	26.6 o	10.00	27.2 0	0.380	0.410	0.33 a	0.370	0.40 D
CNA 5019	26.6 a	9.0 u	27.30	1.28 a	0.300	0.540	0.380	0.390 0.57b
DDA 022022	26.0 a	42.9 a	09.8 C	1.20 a	0.00 a	0.30 a	0.460	0.370
DKA 052055	25.7 a	23.0 C	47.4 d	0.390	0.33 0	0.30 0	0.450	0.370
CNA 0002	55.7 a 24 7 a	17.5 d	39.3 u	0.44 C	0.42 0	0.410	0.380	0.41 D
CNA /000	54.7 a	13.4 u	52.7e	0.50 C	0.55 0	0.580	0.50 0	0.370
AB 002138	34.3 a	27.8 D	59.0 c	0.90 b	0.57 a	0.51 a	0.56 a	0.03 a
Guarani	34.2 a 33 6 a	19.0 d 17 9 d	363 e	0.31c 0.37 c	0.49 b 0.46 b	0.32 a 0.38 h	0.34 a 0.34 h	0.37 b 0.42 h
Mean a ⁽¹⁾	40.8	26.4	48.9	0.77	0.52	0.49	0.50	0.59
Douradão	31.8 h	18.7 d	38.3 P	0.64 c	0.34 h	0.45	0.37 h	0.35
	31.00 31.1 h	13.7 d	29.5 e	0.04 c	0.34 b	0.30 b	0.37 b	0.40 b
CNA 7451	31.10 31.0 h	22.9 c	27.5 C	0.470	0.42 b	0.37 b	0.31 b	0.47 U
CT11801 2 2 2 M	21.0 b	12.9C	22.7 0	0.52 c	0.200	0.32 b	0.300	0.390
4 Massa Propas	20.0 h	12.0 u 18 5 d	22.7 e	0.42 C	0.28 D	0.27 b	0.310	0.29 U
CNA 6697	29.00 28.0h	10.5 u	33.2 e	0.480	0.400	0.44 D	0.270	0.550
CNA 0007	20.00 27.0 h	21.4 C	43.7 d	0.52 C	0.42 D	0.44 D	0.41 D	0.540
UNA 4120	27.00 26.0 h	10.0 U	41.0 d	0.54 C	0.470	0.44 D	0.47 b	0.310
IAC 25	20.90	10.2 d	30.9 e	0.53 0	0.40 0	0.44 0	0.470	0.410
Amor Drange Drange	20.5 D	19.0 d	39.3 U	0.54 C	0.48 0	0.33 D	0.390	0.34 0
AITOZ DIAIICO PIECOCE	20.2 D	19.5 d	41.0 d	0.30 c	0.30 0	0.46 0	0.400	0.480
IKEM 000	25.9 D	19.0 d	44.5 d	0.48 c	0.32 D	0.43 D	0.51 0	0.41 b
BKA 052051	25.0 D	15.2 d	40.9 d	0.40 c	0.34 D	0.41 b	0.37 b	0.32 0
AB 062048	24.7 b	16.1 d	39.7 d	0.44 c	0.35 b	0.29 b	0.276	0.376
BRS Pepita	24.5 D	19.9 d	48.6 d	0.38 c	0.26 b	0.49 a	0.476	0.50 b
CNA /864	24.4 b	20.2 d	38.1 e	0.52 c	0.35 b	0.40 b	0.376	0.476
CT11632-3-3-M	23.3 b	38.8 a	82.2 b	1.51 a	0.85 a	0.56 a	0.47b	0.53 b
BKA 052053	22.6 b	15.4 d	44.0 d	0.65 c	0.30 b	0.29 b	0.34 b	0.36 b
BKA 052034	22.1 b	25.0 c	4/./ d	0./1 c	0.47 b	0.41 b	0.37 b	0.45 b
Noventinha	20.8 b	19.4 d	38.2 e	0.57 c	0.34 b	0.42 b	0.44 b	0.33 b
BRA 01600	19.9 b	26.4 c	50.1 d	0.56 c	0.45 b	0.64 a	0.45 b	0.32 b
BRA 01596	12.4 b	21.4 c	40.1 d	0.62 c	0.52 b	0.66 a	0.39 b	0.37 b
Farroupilha	9.6 b	37.4 a	99.5 a	1.13 b	0.51 b	0.42 b	0.69 a	0.66 a
Mean b ⁽²⁾	24.7	20.8	44.5	0.60	0.41	0.42	0.41	0.42
General mean	33.4	23.8	46.9	0.69	0.47	0.46	0.46	0.51

Table 2: Grain yield (GY), leaf biomass (LeafBio), stem biomass (StemBio) and root density in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth of rice genotypes under water deficit

Means followed by the same letters in the columns do not differ by Scott-Knott' test at 0.05 probability level.

 $\ensuremath{^{(1)}}$ Means of the most productive group according to Scott-Knott' test.

 $^{\scriptscriptstyle (2)}$ Means of the least productive group according to Scott-Knott' test.

C t	GY	LeafBio	StemBio		Root	density (mg	cm ⁻³)	
Genotype -	(g	g per colum	n)	0-0.2 m	0.2-0.4 m	0.4-0.6 m	0.6-0.8 m	0.8-1.0 m
CNA 6187	119.7 a	35.6 b	75.0 b	0.91 b	0.53 c	0.57 a	0.77 a	0.80 b
BRA 052045	118.4 a	38.5 b	65.1 c	1.41 a	0.70 b	0.65 a	0.67 a	0.56 c
Guapa	112.9 a	39.3 b	75.4 b	0.89 b	0.61 b	0.56 a	0.74 a	0.60 c
AB 062138	112.3 a	33.9 b	61.5 c	0.97 b	0.42 c	0.55 a	0.66 a	0.83 b
Arroz Mato Grosso	112.1 a	39.1 b	73.9 b	0.88 b	0.86 a	0.88 a	0.81 a	0.87 b
CNA 4098	111.6 a	32.1 b	56.8 c	0.53 c	0.50 c	0.33 b	0.61 a	0.80 b
BRA 02601	110.5 a	39.2 b	81.1 b	1.15 b	0.56 b	0.82 a	0.99 a	1.02 b
Rio Paranaíba	109.9 a	42.3 a	69.5 b	1.10 b	0.61 b	0.73 a	0.73 a	0.71 c
AB 062041	108.5 a	28.3 c	59.4 c	0.79 c	0.65 b	0.64 a	0.61 a	0.61 c
AB 062037	107.7 a	32.3 b	63.7 c	1.03 b	0.58 b	0.55 a	0.60 a	0.61 c
BRA 052034	107.5 a	31.6 c	59.7 c	0.60 c	0.39 c	0.46 b	0.40 b	0.57 c
Guaporé	106.1 a	43.4 a	59.9 c	0.87 b	0.69 b	0.59 a	0.77 a	0.93 b
CNA 4140	102.9 a	36.7 b	68.6 b	1.01 b	1.03 a	0.67 a	0.82 a	0.79 b
BRA 01600	102.3 a	31.2 c	68.6 b	1.01 b	0.49 c	0.58 a	0.64 a	0.69 c
Urucuí	102.2 a	26.6 c	58.3 c	0.68 c	0.44 c	0.36 b	0.51 b	0.64 c
CT 11632-3-3-M	98.4 a	43.2 a	86.1 b	1.65 a	1.09 a	0.87 a	0.73 a	0.92 b
BRA 032051	97.8 a	24.1 d	66.5 c	0.75 c	0.45 c	0.47 b	0.69 a	0.68 c
BRA 052023	96.3 a	29.4 c	65.2 c	0.56 c	0.77 b	0.52 a	0.48 b	0.64 c
CNA 5018	96.1 a	45.5 a	83.6 b	1.41 a	0.53 c	0.72 a	0.68 a	0.88 b
BRS Esmeralda	95.7 a	31.1 c	57.5 c	0.63 c	0.68 b	0.70 a	0.61 a	0.82 b
CNA 4128	95.6 a	24.9 d	59.3 c	0.95 b	0.88 a	0.67 a	0.75 a	0.91 b
BRA 052053	95.1 a	24.6 d	59.8 c	0.85 b	0.39 c	0.46 b	0.49 b	0.57 c
CNA 4137	93.6 a	21.1 d	50.1 d	0.55 c	0.60 b	0.54 a	0.59 a	0.62 c
BRA 032033	91.1 a	27.7 с	66.7 c	0.71 c	0.62 b	0.70 a	0.90 a	1.03 b
BRS Pepita	91.0 a	25.8 c	55.8 c	0.62 c	0.35 c	0.52 a	0.60 a	0.74 c
CNA 7864	90.8 a	25.1 d	56.9 c	0.87 b	0.74 b	0.50 a	0.62 a	0.66 c
IAPAR 9	90.1 a	24.2 d	50.8 d	0.70 c	0.55 c	0.51 a	0.57 a	0.58 c
BRSMG Caravera	89.6 a	23.4 d	52.3 d	0.55 c	0.45 c	0.27 b	0.39 b	0.56 c
Mean a ⁽¹⁾	102.3	32.1	64.5	0.88	0.61	0.58	0.66	0.74
AB 062048	88.2 b	20.5 d	48.9 d	0.44 c	0.38 c	0.36 b	0.33 b	0.38 c
TOX 503-4-115-B-B	87.7 b	37.7 b	76.9 b	1.29 a	0.61 b	0.67 a	0.70 a	0.83 b
BRA 01596	87.2 b	24.1 d	42.3 d	0.37 c	0.29 c	0.19 b	0.22 b	0.32 c
CNA 6687	85.1 b	26.7 c	57.3 c	0.70 c	0.50 c	0.37 b	0.46 b	0.72 c
CNA 7680	83.0 b	25.9 с	62.0 c	0.74 c	0.45 c	0.51 a	0.69 a	0.99 b
CNA 7451	81.7 b	24.4 d	64.9 c	0.59 c	0.54 c	0.46 b	0.60 a	0.65 c
Douradão	81.2 b	27.3 c	53.9 d	0.48 c	0.28 c	0.28 b	0.42 b	1.48 a
4 Meses Branco	80.3 b	22.0 d	52.2 d	0.54 c	0.43 c	0.32 b	0.47 b	0.47 c
Guarani	80.0 b	21.5 d	53.3 d	0.70 c	0.59 b	0.57 a	0.58 a	0.85 b
Arroz Branco Precoce	78.8 b	24.1 d	53.1 d	0.76 c	0.63 b	0.54 a	0.64 a	0.75 b
BRS Bonança	78.4 b	26.9 c	52.1 d	1.15 b	0.58 b	0.48 b	0.65 a	0.90 b
CNA 5166	77.3 b	23.7 d	50.2 d	0.41 b	0.40 c	0.32 b	0.36 b	0.54 c
Mean b ⁽²⁾	82.4	25.4	55.6	0.68	0.47	0.42	0.51	0.74
IREM 247	71.1 c	21.4 d	55.8 c	0.44 c	0.40 c	0.38 b	0.51 b	0.50 c
CNA 6682	68.9 c	26.5 c	59.5 c	0.61 c	0.36 c	0.36 b	0.45 b	0.49 c
Farroupilha	68.9 c	37.6 b	106.9 a	1.31 a	0.60 b	0.60 a	0.48 b	0.85 b
CT 11891-3-3-3-M	67.5 c	19.1 d	38.5 d	0.36 c	0.26 c	0.29 b	0.32 b	0.46 c
IAC 25	67.2 c	24.7 d	56.7 c	0.68 c	0.44 c	0.38 b	0.52 b	0.59 c
IREM 656	64.1 c	27.2 c	53.5 d	0.97 b	0.52 c	0.53 a	0.72 a	0.91 b
BRSMG 355	58.6 c	22.2 d	53.0 d	0.52 c	0.36 c	0.41 b	0.58 a	0.98 b
Noventinha	49.2 c	23.7 d	44.6 d	0.39 c	0.23 c	0.28 b	0.31 b	0.45 c
Mean c ⁽³⁾	64.4	25.3	58.6	0.66	0.40	0.40	0.49	0.65
General mean	91.0	29.3	61.3	0.79	0.54	0.51	0.59	0.72

Table 3: Grain yield (GY), leaf biomass (LeafBio), stem biomass (StemBio) and root density in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth of well-watered rice genotypes

Means followed by the same letters in the columns do not differ by Scott-Knott' test at 0.05 probability level.

⁽¹⁾ Means of the most productive group according to Scott-Knott' test.

⁽²⁾ Means of the intermediate productive group according to Scott-Knott' test.

⁽³⁾ Means of the least productive group according to Scott-Knott' test.

with the lowest P content and 0.45 mg cm⁻³ with the P content of 200 mg dm⁻³. Since, for the same P treatments, the shoot biomass (leaves + stems) was 69.2 and 72.3 g, the root:shoot ratio was higher for the lowest P content, corroborating the previous authors.

Since the genotypes were tolerant to P-deficient soils, the water deficit had more impact on grain yield than the P supply. As the water deficit was imposed at panicle emission stage, and root weight density peaks at around flowering (Kato & Okami, 2010), the higher root density, although not significant, may have contributed to minimize the effect of water deficit and explain the highest grain yield at the lowest P content. These results corroborate the association between the tolerance to P deficiency and adaptation to abiotic stresses (Chin *et al.*, 2010; 2011).

Regarding the well-watered treatment, as the genotypes did not suffer water deficit, the increase in root density as a response to P deficiency is less important to define the grain yield than the P supply. The plant phosphorus uptake is related to the diffusive flux of phosphorus in soil and this is influenced by the soil moisture and P concentration in the soil (Costa *et al.*, 2006).

The genotypes differed significantly for all the agronomic traits. Furthermore, it was observed that water regimes influenced differently grain yield, stem biomass and root density of genotypes in all layers evaluated except in the surface layer, as the water regime x genotype interaction was significant for these agronomic traits (Table 1).

Centrito *et al.* (2009), applying water deficit from the flowering stage until the beginning of physiological maturity, also found a significant decrease in aboveground dry mass and grain yield, that varied among rice genotypes, being the most productive those genotypes which showed higher photosynthesis and mesophyll and stomata conductances. Kamoshita *et al.* (2004) and Guimarães *et al.* (2013; 2016) also observed genetic variation in rice performance when subjected to water deficit.

Differences among genotypes for the traits evaluated, in both water regimes, can be observed in Tables 2 and 3. As there was an interaction between water regime and genotype, probably the root density is not a constitutive trait for all genotypes. The root development as a constitutive trait gives advance preparation for the plant to support possible occurrences of water deficit. Considering that the water deficit periods are unpredictable, both in intensity, durability and time of occurrence, this, along with others, is an essential characteristic of cultivars recommended for areas with the occurrence of uneven rainfall distribution.

Ji *et al.* (2012) found that, for a sensitive rice cultivar, a 20-days drought stress at reproductive stage caused

reduction of 61% and 14% in root mass from 0 to 0.3 m and from 0.3 to 0.9 m, respectively. However, for a tolerant rice cultivar it was observed a reduction of 43% in root mass from 0 to 0.3 m but an increase of 72% from 0.3 to 0.9 m.

Scott-Knott's test classified the genotypes in two and three groups in relation to grain yield under water deficit and under well-watered conditions, respectively (Tables 2 and 3). The most productive group under water deficit, with 54.2% of the genotypes evaluated, showed the following means: 40.8 g per column of grain yield, 26.4 g per column of leaf biomass, 48.9 g per column of stem biomass and root density in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth of 0.77, 0.52, 0.49, 0.50 and 0.59 mg cm⁻³, respectively (Table 2). Under wellwatered conditions, the most productive group showed the following means: 102.3 g per column of grain yield, 32.1 g per column of leaf biomass, 64.5 g per column of stem biomass and root density in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth of 0.88, 0.61, 0.58, 0.66 and 0.74 mg cm⁻³, respectively (Table 3).

Grain yield under water deficit correlated positively and significantly with the root density in the deeper layers of the soil (Table 4). This was due, probably, to the biggest water uptake afforded by a deeper root system, which favored the accumulation of biomass in the leaves and stems. Matsuo *et al.* (2009), evaluating the performance of rice genotypes under water deficit conditions, observed a correlation between the water uptake capacity and the accumulation of biomass in shoots.

The leaf and stem biomass were positively correlated with grain yield and root density in all soil layers. Presumably, plants with higher leaf biomass also show greater leaf area, therefore, greater photosynthesis ability, which would lead to a greater accumulation of carbohydrates in the stems.

Among the 26 genotypes classified in the most productive group under water deficit, the genotypes AB 062037, AB 062041, AB 062138, Arroz Mato Grosso, BRA 02601, BRA 032033, BRA 052023, BRA 052045, BRS Esmeralda, BRSMG Caravera, CNA 4098, CNA 4137, CNA 4140, CNA 5018, CNA 6187, Guapa, Guaporé, Rio Paranaíba and Urucuí were also classified in the most productive group under well-watered conditions (Tables 2 and 3). With few exceptions, this group of 26 genotypes is that with high root density in the deepest layers of soil (Table 2), which is in accordance with the result of the linear correlation analysis.

The use of water by genotypes were evaluated considering their classification in clusters, according to the Scott-Knott's test. In the two water regimes, the cumulative evapotranspiration of soil columns was higher for those with genotypes classified in the most productive group (Figure 1). Under water deficit conditions, the soil

Table 4: Pearson's correlation coefficient among means of grain yield (GY), leaf biomass (LeafBio), stem biomass (StemBio) and root density in the soil layers of 0-0.2, 0.2-0.4, 0.4-0.6, 0.6-0.8 and 0.8-1.0 m depth, under water deficit. n = 48

Variable	LeafBio		StemBio —	Root density						
				0-0.2 m	0.2-0.4 m	0.4-0.6 m	0.6-0.8 m	0.8-1.0 m		
GY	0.401 *	0.383 *	0.296 ns	0.228 ns	0.432 *	0.539 **	0.443 *			
LeafBio			0.922 **	0.822 **	0.716 **	0.608 **	0.724 **	0.773 **		
StemBio				0.785 **	0.565 **	0.584 **	0.719 **	0.692 **		
Root density	0-0.2 m				0.727 **	0.525 **	0.618 **	0.739 **		
	0.2-0.4 m					0.645 **	0.593 **	0.719 **		
	0.4-0.6 m						0.707 **	0.647 **		
	0.6-0.8 m							0.854 **		

ns, * and **: not significant, significant at 0.05 and 0.01 levels by t-test, respectively.



Figure 1: Cumulative evapotranspiration of soil columns cropped with upland rice genotypes, from panicle emission to harvest, according to statistic clusters. A. W/ ws and W/o ws – with and without water deficit, respectively, Ga and Gb – the most and the least productive group, respectively. B. Ga, Gb, Gc - groups in descending order of grain yield under well-watered conditions.

columns with the genotypes of the most productive group showed on average cumulative evapotranspiration of 30.1 L per column and those of the least productive group 27.0 L per column, for a similar period of 44 days after panicle emission. The three clusters defined for the well-watered conditions, in descending order of grain yield, showed cumulative evapotranspiration of 62.1, 52.5 and 47.8 L per column, respectively.

As the plants totally covered the soil in the columns at the panicle emission stage, transpiration accounted for most of the water lost to the atmosphere from this stage to harvest. According to Blum (2009), since biomass production is tightly linked to transpiration, breeding for maximized soil moisture capture for transpiration is the most important target for yield improvement under water deficit. Zain *et al.* (2014) found that 15-day water deficit cycle reduced rice transpiration rate by 42% and, as it has a positive correlation with net photosynthesis rate, reduced grain yield.

Greater root length density increases the water storage capacity of the root zone, and a deeper root system is associated with more water uptake from the soil and with

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better crop performance under drought conditions (Mishra & Salokhe, 2011). Kato & Okami (2010) noted that when the soil water potential at 20-cm depth was below -50 kPa the stomatal behavior reflected the rice root growth in the subsurface layer. These results suggest the role of vigorous root growth in soil water uptake and hence, in maintaining transpiration in upland rice crop.

CONCLUSIONS

The water deficit had more impact on grain yield than the P supply.

The genotypes AB 062037, AB 062041, AB 062138, Arroz Mato Grosso, BRA 02601, BRA 052045, CNA 4098, CNA 6187, Guapa, Guaporé and Rio Paranaíba were classified in the most productive group under both water regimes.

The most productive genotypes under water deficit showed higher root density in the deeper soil layers.

The most productive genotypes in the two water regimes were also those that showed the highest transpiration.

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