COLD TOLERANCE AT THE GERMINATION STAGE OF RICE: METHODS OF EVALUATION AND CHARACTERIZATION OF GENOTYPES

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ABSTRACT: Rice cold tolerance at the germination stage is important in Rio Grande do Sul (RS) where temperatures below 15°C prevent or reduce germination and plant establishment in early sowings. The present study aimed at identifying an adequate method for cold tolerance evaluation of the rice germination stage and at verifying the variability among 24 rice genotypes of different origins. Cold tolerance was evaluated in experiment I, germination under two conditions: 13°C for 28 days and 28°C for seven days, and in experiment II, germination under 28°C for 72 hours, 13°C for 96 hours and again 28°C for 72 hours. In experiment I measured characteristics were germination index, percentage of seeds with coleoptile length superior to 5 mm and percentage of reduction in coleoptile length due to cold. In experiment II the measured characteristic was coleoptile regrowth after the cold period. Cold tolerance varied among genotypes studied in both experiments, but only the percentage of reduction in coleoptile length and coleoptile regrowth allowed a better distinction between the tolerant checks and the susceptible one. In general, genotypes belonging to the Japonica subspecies presented higher cold tolerance than Indica, but there was variability within subspecies. The most adequate method of evaluation of cold tolerance is through percentage of reduction in coleoptile length and coleoptile regrowth. Among Japonica genotypes, Quilla 64117 and Diamante presented the highest cold tolerance, and among Indica, cultivars BR-IRGA 410 and IRGA 416 were the most cold tolerant at the germination stage.

Key words: Oryza sativa L., low temperature, variability, Indica and Japonica subspecies

TOLERÂNCIA AO FRIO NO ESTÁDIO DE GERMINAÇÃO EM ARROZ: MÉTODOS DE AVALIAÇÃO E CARACTERIZAÇÃO DE GENÓTIPOS

RESUMO: A tolerância ao frio em arroz no estádio de germinação é importante no Rio Grande do Sul (RS) onde temperaturas abaixo de 15°C impedem ou reduzem a germinação e o estabelecimento das plantas em semeaduras precoces. O presente trabalho teve por objetivos identificar uma metodologia adequada para avaliação da tolerância ao frio no final da germinação em arroz e verificar a variabilidade existente entre 24 genótipos de arroz de diferentes origens. A tolerância ao frio foi avaliada em dois experimentos: experimento I, com germinação sob duas condições: 13°C por 28 dias e 28°C por sete dias, e experimento II, com germinação a 28°C por 72 horas, 13°C por 96 horas e novamente a 28°C por 72 horas. No experimento I as características medidas foram o índice de germinação, porcentagem de sementes com coleóptilo superior a 5 mm e porcentagem de redução no comprimento do coleóptilo devido ao frio. Na experimento II a característica medida foi o recrescimento do coleóptilo após o período de frio. Houve variabilidade para tolerância ao frio entre os genótipos estudados em ambos os experimentos, mas apenas a porcentagem de redução no comprimento do coleóptilo e o recrescimento do mesmo permitiram melhor distinção entre as testemunhas tolerantes e a sensível. Os genótipos da subespécie Japônica apresentaram maior tolerância ao frio que os Indica, no entanto, houve variabilidade dentro das subespécies. O método mais adequado de avaliação da tolerância ao frio é pela porcentagem de redução no comprimento do coleóptilo e recrescimento do mesmo. Entre os genótipos Japônica, Quilla 64117 e Diamante, apresentam maior tolerância ao frio, e entre os Indica, as cultivares BR-IRGA 410 e IRGA 416 são as mais tolerantes ao frio no estádio de germinação.

Palavras-chave: Oryza sativa L., baixa temperatura, variabilidade, subespécies Indica e Japônica

INTRODUCTION

In Rio Grande do Sul, Brazil, rice is sown from September to December, according to the region and climatic conditions. The sowing time recommended for most medium-cycle cultivars, in the majority of the regions is, however, concentrated from early October to middle November, a period with average temperatures around 15°C.
(Oliveira, 1997). Even though temperature does not prevent rice germination, it delays beginning and, consequently, plant emergence. Optimum temperature range for rice germination lies between 20 and 35°C, and the temperature of 10°C is cited as the minimum critical value below which rice does not germinate (Yoshida, 1981b).

Good performance during germination is important to guarantee fast establishment and uniform crop stand (Krishnasamy & Seshu, 1989). In Rio Grande do Sul, most rice cultivars belong to Indica subspecies, which present slow and not uniform germination under cold temperature, resulting in irregular emergence and low plant population (Souza, 1990). Since management practices can not minimize the problem, genetic breeding for cold tolerance at the germination stage may contribute to a better crop establishment in early sowings.

Selection of cold tolerant genotypes under field conditions is employed in countries such as Korea (Heu & Bae, 1972) and United States (Carnahan et al., 1972), but is not efficient in Rio Grande do Sul because of the weather instability. Relatively mild, short winter seasons do not present good selection pressure. Rice cold tolerance at the germination stage has been studied under controlled temperature conditions, which allows adjusting intensity and duration, as well as greater precision resulting from the absence of other abiotic and biotic factors that may interfere in field data (Blum, 1988). According to methods of evaluation available in the literature, seeds are submitted to temperatures varying from 10 to 25°C for periods of three to thirty five days and characteristics most commonly measured are germination percentage and speed, and coleoptile and radicule length (Maya, 1988; Srinivasulu & Vergara, 1988; Bertin et al., 1996; Sthapit & Witcombe, 1998).

These methodologies are important mainly for cold tolerance inheritance studies, in which stress control is fundamental to obtain precise results. For rice, Sthapit & Witcombe (1998) reported heritability varying from 0.74 to 0.87 for percentage and index of germination, evaluated after submitting seeds to 17°C for seven days. Ratho & Pradhan (1991) identified a male sterile line with high cold tolerance at the germination stage, which has been pointed out as having cytoplasmatic inheritance. For maize, an inheritance study of cold tolerance at the germination stage was also performed under controlled temperature and allowed the identification of additive-dominant gene action for emergence percentage and initial growth (Revilla et al., 2000).

However, most studies performed under controlled temperature at the rice germination stage are related to the identification of variability and genotype characterization. They were responsible for identifying genotypes of better performance during germination under cold temperatures and, in general, it is known that genotypes belonging to Japonica subspecies are more cold tolerant than Indica (Li et al., 1981; Cruz & Milach, 1999). Genetic variability for cold tolerance among wild rice lines was also studied (Suh et al., 1997), and three QTLs (quantitative trait loci) related to low temperature germinability were mapped for a population resulting from a cross between a wild and a cultivated line (Suh et al., 1999).

Currently, however, there is a lack of information concerning the behavior of Brazilian rice genotypes in relation to their germination under controlled cold temperatures, and indication of the most adequate methodology for application in breeding programs. In this context, the present study aimed at establishing a methodology of evaluation of cold tolerance at the germination stage under controlled temperature, aiming to identify the most adequate characteristics to be used as evaluation criteria, and to characterize the variability among 24 rice genotypes from the germplasm bank of the Instituto Rio Grandense do Arroz (IRGA) regarding to these characteristics.

MATERIAL AND METHODS

Twenty four rice genotypes were studied, 12 belonging to Japonica and 12 to Indica subspecies (Table 1). They were chosen because they have different origins and represent a wide range of variability within the rice germplasm. Six of these genotypes are checks (five tolerant and one sensitive) which were included based on observation of their field performance under cold weather. Among the Indica genotypes, six are rice cultivars recommended for planting in RS (BR-IRGA 409, BR-IRGA 410, IRGA 416, IRGA 417, BR 7 – Taim, and El Paso 144).

Two methods were used to characterize genotypes cold tolerance, described in Experiments I and II. In Experiment I, seeds of 24 rice genotypes were germinated under two conditions: 13°C for 28 days (cold) and 28°C for seven days (control). For each genotype, seeds from the germplasm bank of IRGA, harvested in the same cropping season, were selected based on size uniformity and absence of spots. They were sterilized with ethanol 70% for 30 seconds and sodium hipoclorite 5% for 20 minutes, and washed six times with sterile distilled water. Seeds were placed on Petri dishes containing two layers of germination paper, wet with distilled water and 1 mL of Benomyl solution 2.5 ppm to avoid contamination. The experiment was conducted in a randomized block design with three replications, and the blocks constituted of different shelves in the germination chamber. Each Petri dish contained 20 seeds, and the average of these seeds was used as a replication, amounting to 60 seeds per genotype.

Seeds germinated at 13°C had their coleoptile length measured weekly for a period of 28 days and for seeds germinated at 28°C this measure was made seven days after the beginning of the experiment. Evaluation of
Rice cold tolerance at the germination stage

The genotypes cold tolerance in this experiment was carried out by means of the following characteristics:

• Germination index (GI): \( GI = \frac{N_{14} + N_{21}}{20} \times 100 \), where \( N_{14} \) = number of germinated seeds 14 days after the beginning of the cold treatment; \( N_{21} \) = number of germinated seeds 21 days after the beginning of the cold treatment; 20 being the total number of seeds per genotype per replication. For the GI calculation only the seeds presenting coleoptile and radicule were considered.

• Percentage of seeds with coleoptile superior to 5 mm (PERCOL5): obtained considering all the germinated seeds 28 days after the beginning of the cold treatment and by verification of the percentage that presented coleoptile length superior to 5 mm, according to the formula:

\[
\text{PERCOL5} = \frac{\text{number of seeds with coleoptile > 5 mm}}{20} \times 100
\]

• Percentage of reduction in coleoptile length (REDCOL): obtained through comparison of average coleoptile length 28 days after germination at 13°C (cold treatment) with that obtained 7 days after germination at 28°C (control), and calculating the percentage of reduction in coleoptile length by germination under cold temperature, according to the formula:

\[
\text{REDCOL} = \left[ \frac{\text{col. length under cold temperature} \times 100}{\text{col. length under control}} \right] - 100
\]

In Experiment II, seeds of the 24 rice genotypes were submitted to germination under the following conditions: 28°C for 72 hours, 13°C for 96 hours and again 28°C for 72 hours. This procedure was used to simulate field conditions, where temperature variation is ordinarily observed. The choice and sterilization of the seeds and conduction of the experiment were made according to Experiment I. Coleoptile length was measured in two occasions: after the period of 96 hours at 13°C (LENGTH 1) and after the second period of 72 hours at 28°C (LENGTH 2). Cold tolerance evaluation was performed through coleoptile regrowth (COLREG; mm), which consisted in the difference between the second and the first measurements, or in how much the coleoptile grew under normal temperature after the cold treatment, according to the formula: 

\[
\text{COLREG} = (\text{LENGTH 2}) - (\text{LENGTH 1})
\]

Both experiments were conducted at 13°C because this temperature was the one used by Maya (1988) to evaluate cold tolerance at the germination stage of the checks employed in this study. Data from both experiments were submitted to residual analysis to verify normality and variance homogeneity, and to regression analysis to test if data transformation was necessary. The latter indicated the need to transform germination index (Experiment I) through the square root, and coleoptile regrowth (Experiment II) through logarithm. The two other characteristics, percentage of seeds with coleoptile superior to 5 mm and percentage of reduction in coleoptile length, presented a normal distribution and were not transformed.

Data were, then, submitted to analysis of variance and means comparison by the Tukey test \( (P = 0.05) \). Relationships among the measured characteristics were established through Pearson’s coefficient correlation. All analysis were performed in the Statistical Analysis System (SAS Institute, 2000).

RESULTS AND DISCUSSION

There was variability for cold tolerance at the germination stage among the genotypes (Table 2). The percentage of reduction in coleoptile length (REDCOL) pre-
presented the highest coefficient of variation, probably because this characteristic was not transformed and also because it was the only characteristic obtained through comparison of the germination capacity between two contrasting conditions (cold and control), which involved different sets of seeds for a same genotype (treatment).

In relation to the germination index (GI), that expresses the germination speed under cold temperature, IRGA 417 presented values superior to 80%, differing significantly from the others (Figure 1). These results indicate that more seeds of this genotype had germinated 14 and 21 days after the beginning of the experiment. Germination speed is important for crop establishment; however, it does not necessarily hold any relation with the ability of a genotype to elongate coleoptile and radicle under cold temperature. As a matter of fact, for the GI calculation only seeds that presented coleoptile and radicle were considered, independent of their length. Germination speed, and consequently GI, are related to a high seed vigour and this may be the cause of the better performance of IRGA 417 for GI.

Percentage of seeds with coleoptile length greater than 5 mm (PERCOL5) after 28 days of germination at 13°C, by its turn, takes into consideration coleoptile length, and despite the wide variation observed among the genotypes, there was no clear separation between genotypes (Figure 2). Fourteen genotypes presented more than 70% of the seeds with coleoptile length superior to 5 mm, including the tolerant checks (Koshihikari, Quilla 64117, Diamante, Caloro and Quilla 66304), and the cold sensitive IR 8. This characteristic reflected the coleoptile elongation capacity under cold temperature, and both IR 8 and IRGA 417, superior for GI, also presented high capacity of coleoptile growth, with values greater than 90% for PERCOL5 (Figure 2). This characteristic is a good criterion to distinguish cold-tolerant from cold-sensitive rice genotypes (Maya, 1988). In fact the five genotypes used as cold tolerance parameters presented high percentage of seeds with coleoptiles longer than 5 mm, the same being true for the sensitive check IR 8 (Figure 2). What demonstrates that this characteristic did not tell apart tolerant and sensitive checks in this case.

Percentage of reduction in coleoptile length by cold (REDCOL) varied from 11 to 68%, and although there was no significant difference, the cold tolerant checks presented lower reductions in coleoptile length due to cold when compared to the sensitive IR 8 (Figure 3). All genotypes with reduction smaller than 25% belonged to Japonica subspecies, with exception of BR-IRGA 410, which is an Indica genotype. The lower values for REDCOL were observed for the tolerant checks Diamante, Koshihikari and Quilla 64117 (Figure 3). This characteristic expresses the amount of reduction in genotypes coleoptile length by germination under cold in relation to normal temperature. In this way, the effect of vigour differences among seed lots are minimized.

Table 2 - Mean squares, coefficients of variation and coefficients of determination for the four evaluated characteristics.

<table>
<thead>
<tr>
<th>Sources of variation</th>
<th>GI</th>
<th>PERCOL5</th>
<th>REDCOL</th>
<th>COLREG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genotype</td>
<td>7.98**</td>
<td>1734.7**</td>
<td>776.3**</td>
<td>0.7**</td>
</tr>
<tr>
<td>Blocks</td>
<td>0.15</td>
<td>64.1</td>
<td>375.4</td>
<td>0.2</td>
</tr>
<tr>
<td>Error</td>
<td>0.35</td>
<td>215.3</td>
<td>97.7</td>
<td>0.1</td>
</tr>
<tr>
<td>CV (%)</td>
<td>10.4</td>
<td>20.7</td>
<td>29.0</td>
<td>13.0</td>
</tr>
<tr>
<td>R²</td>
<td>0.92</td>
<td>0.80</td>
<td>0.81</td>
<td>0.80</td>
</tr>
</tbody>
</table>

** Significant at 1%.

GI = germination index, PERCOL5 = percentage of seeds with coleoptile length greater than 5 mm, REDCOL = percentage of reduction in coleoptile length, COLREG = coleoptile regrowth growth

Sci. Agric. (Piracicaba, Braz.), v.61, n.1, p.1-8, Jan./Fev. 2004
Data relative to coleoptile regrowth (COLREG) yielded values varying from 4.3 mm to 28.3 mm (Figure 4), indicating that the germination process after the cold period was differently affected among genotypes. Three tolerant checks (Diamante, Quilla 66304 and Quilla 64117) formed a group with high coleoptile regrowth, superior to 25 mm. The two other cold tolerant checks, Caloro and Koshihikari, presented intermediate regrowth as well as the sensitive check, IR 8 (Figure 4).

One of the objectives of the present study was to identify a methodology capable of differentiating cold tolerant from cold sensitive genotypes, at the germination stage. Among the characteristics studied, REDCOL was the one that allowed for a better distinction between the tolerant and sensitive checks (Figure 3). Comparison of growth under cold temperature in relation to normal temperature must present a better estimation of a genotype cold answer and this was exactly what was achieved through the percentage of reduction in coleoptile length in this study (Miedema, 1982). In the case of GI and PERCOL5, however, it was not possible to distinguish the tolerant from cold sensitive genotypes, at the germination stage. Among the characteristics studied, REDCOL was one of the methodologies that allowed for a better distinction between the tolerant and sensitive checks (Figure 3). Comparison of growth under cold temperature in relation to normal temperature must present a better estimation of a genotype cold answer and this was exactly what was achieved through the percentage of reduction in coleoptile length (REDCOL) and coleoptile regrowth (COLREG) evaluated in 24 rice genotypes.

![Figure 3 - Percentage of reduction in coleoptile length of 24 rice genotypes](image)

**Figure 3** - Percentage of reduction in coleoptile length of 24 rice genotypes, obtained through comparison of their germination under cold temperature (13°C for 28 days) with their germination under normal temperature (28°C for seven days), compared by the Tukey test (α=0.05).

![Figure 4 - Coleoptile regrowth (mm) of 24 rice genotypes](image)

**Figure 4** - Coleoptile regrowth (mm) of 24 rice genotypes submitted to germination at 28°C for 72 h, 13°C for 96 h and 28°C for 72 h, compared by the Tukey test (α=0.05).

**Table 3** - Correlation coefficients between the characteristics, germination index (GI), percentage of seeds with coleoptile greater than 5 mm (PERCOL5), percentage of reduction in coleoptile length (REDCOL) and coleoptile regrowth (COLREG) evaluated in 24 rice genotypes.

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>GI</th>
<th>PERCOL5</th>
<th>REDCOL</th>
<th>COLREG</th>
</tr>
</thead>
<tbody>
<tr>
<td>GI</td>
<td>1.0</td>
<td>0.7**</td>
<td>-0.3**</td>
<td>0.2</td>
</tr>
<tr>
<td>PERCOL5</td>
<td></td>
<td>1.0</td>
<td>-0.6**</td>
<td>0.2</td>
</tr>
<tr>
<td>REDCOL</td>
<td></td>
<td></td>
<td>-0.3**</td>
<td></td>
</tr>
<tr>
<td>COLREG</td>
<td></td>
<td></td>
<td></td>
<td>1.0</td>
</tr>
</tbody>
</table>

**Table 4** - Proposed criteria for characterization of the rice genotypes studied as to their cold tolerance at the germination stage evaluated by means of the percentage of reduction in coleoptile length (REDCOL) and coleoptile regrowth (COLREG).

<table>
<thead>
<tr>
<th>Cold reaction</th>
<th>REDCOL</th>
<th>COLREG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Highly tolerant</td>
<td>equal or less than 25</td>
<td>equal or greater than 20</td>
</tr>
<tr>
<td>Tolerant</td>
<td>equal or less than 25</td>
<td>between 8 and 20</td>
</tr>
<tr>
<td>Intermediate</td>
<td>between 25 and 50</td>
<td>equal or greater than 50</td>
</tr>
<tr>
<td>Sensitive</td>
<td>equal or greater than 50</td>
<td>equal or greater than 20</td>
</tr>
<tr>
<td>Highly sensitive</td>
<td>equal or greater than 50</td>
<td>equal or less than 8</td>
</tr>
</tbody>
</table>
two tolerant checks from the sensitive IR 8 (Figure 4). In this character, only ten days were enough to obtain the results, and the temperature variation allowed for simulating what happens under field conditions, where periods of cold temperature are inserted between periods of high temperature. In addition, the initial temperature of 28°C aimed at minimizing intrinsic differences of vigour among the genotypes, providing ideal germination conditions to all seeds. Therefore, differences in COLREG among genotypes may be an indicative of a distinct capacity of recovery of the germination process after a cold period, and of different degrees of cold tolerance.

Both REDCOL and COLREG seem to be the most adequate characteristics to be used to evaluate cold tolerance during the germination period in rice. Correlation between them was negative as expected (Table 3), proving that a greater coleoptile regrowth was accompanied by a lower reduction in coleoptile length due to cold temperature. However, this correlation was low, indicating that some genotypes considered tolerant based on a bigger COLREG, may be considered sensitive based on their REDCOL, and vice-versa. It is thus worthy discussing whether the low correlation between the REDCOL and COLREG indicates either they are related to different aspects of cold tolerance or not.

Germination is divided into three phases: imbibition, activation and post-germination growth (Yoshida, 1981a). The largest effects of cold temperature during germination seem to be associated to the imbibition phase, considered the most sensitive (Simon apud Blum, 1988). Cold temperature during this phase leads to increasing escape of solutes from the seeds, such as aminoacids and carbohydrates, which has been attributed to the incomplete plasma membrane of the dry seed and to the disturbance caused on its reconstruction during imbibition phase by cold temperature.

Cellular membrane is considered the main target for cold damage and the primary cause of other metabolic disturbances observed within cells (Lyons, 1973). The reason for this is the phase transition that takes place in the plasma membrane under cold temperature, which changes from a liquid-crystallin state to a solid-gel state, leading to physical contraction and opening of channels that increase its permeability. Differences in the composition of the lipid layer were pointed out as responsible for the greater tolerance to the phase transition under cold temperature, in such a way that the greater the lipid insaturation degree, the larger the cold tolerance, because the phase transition would occur at a lower temperature (Murata & Yamaya, 1984).

Differences in lipid composition were observed between cold tolerant and sensitive rice genotypes, with the first ones presenting a larger proportion of insaturate fatty acids (Majumder et al., 1989). Similarly, Bertin (1998) verified alterations in the fatty acid composition of the lipid layer in rice somaclones presenting higher cold tolerance than the original genotypes. It may thus be speculated that the differential behavior of some genotypes when evaluated through REDCOL and COLREG was related to the imbibition phase of the seeds that occurred at 13°C in the first and at 28°C in the second.

In spite of the imbibition being considered the most sensitive phase, Yoshida (1981a) reports that the greatest influence of temperature on germination occurs in the subsequent phases of activation and growth of coleoptile and radicle. The reduction in coleoptile growth during these phases may be attributed to the direct effect of cold temperature on cellular elongation and division, or to its indirect effect leading to a metabolic unbalance (Lyons, 1973). These phases must have been affected in both characteristics, but, while in REDCOL cold temperature was constant during all 28 days of germination, in COLREG it lasted for only four days. Once again, each characteristic evaluated different aspects, the first case being the capacity of coleoptile to grow under permanent cold and; in the latter, the capacity of growth recovery under normal temperature after cold stress.

Physiological mechanisms involved with cold tolerance at the vegetative period are better understood. Among the many processes that have been studied and published, the regulating role of kinases dependent on calcium and activated by cold temperature exposition (Saijo et al., 2000; Martín & Busconi, 2001); the involvement of abscisic acid as a signalling molecule (Lee et al., 1995); and the levels of activation of the enzymes associated to oxidative stress such as ascorbate peroxidase (Sato et al., 2001), catalase and glutathione reductase (Kang & Saltveit, 2002) may be cited. Despite the lack of knowledge regarding the germination period of rice, the existence of different physiological mechanisms involved with cold tolerance at the vegetative period is an indication that along the germination phases, different processes may also be affected, implying in different tolerance mechanisms. For instance, Massardo et al. (2000) studied two oat cultivars, one tolerant and another sensitive, in relation to the physiological aspects involved during germination under cold temperature (3°C), and demonstrated that higher metabolic rates and less oxidative damage were the answers of the tolerant genotype to cold temperature during germination.

Under these premises, the low correlation between REDCOL and COLREG may be attributed to the fact that they are associated to different cold tolerance mechanisms at the germination period, and yield complementary results in cold tolerance evaluation, that is, genotype characterization for cold tolerance at the germination stage should consider the results of both characteristics simultaneously. Based on the extent of variation obtained for REDCOL and COLREG, and on the perfor-
mance of the tolerant checks in this study, a criterion for characterization of the genotypes, in relation to their cold reaction at the germination stage, is proposed (Table 4). It comprises five classes of cold reaction that are determined by nine different combinations of values for REDCOL and COLREG. Considering this classification, attention shall be drawn to genotypes Quilla 64117 and Diamante, once they are highly cold tolerant at the germination stage (Figures 3 and 4). It was also observed that the genotype IR 8, though used as a sensitive check, was not inferior to the other genotypes, showing an intermediate or even superior behavior to many of them (Figures 3 and 4). Indica genotypes generally need higher temperatures to germinate than Japonica, and the same is true for photoperiod sensitive and insensitive genotypes, indicating a higher sensitivity to cold temperature of Indica and photoperiod sensitive genotypes (Oka apud Takahashi, 1984). This might be an explanation for the intermediate behavior of IR 8, which is a photoperiod insensitive genotype.

The variability among the rice genotypes was expected because of their different origins and the two subpecies they belong to. However, variability for cold tolerance was also observed within each subspecies. The availability of cold tolerance sources within the Indica subspecies is interesting for breeding cold tolerant genotypes to be cultivated in Rio Grande do Sul, since it allows them to be used as genitors in crosses, preventing the sterility problems that result from Indica x Japonica crosses (Kubo & Yoshimura, 1999; Kubo et al., 2000) as well as the transfer of undesirable characters from the Japonica genotypes to the adapted Indica ones. In this sense, the genotype BR-IRGA 410 may be considered a possible cold tolerance source among the Indica genotypes studied, once the values obtained for REDCOL and COLREG (Figures 3 and 4) allow to classify it as a tolerant genotype (Table 4).

In the breeding process for cold tolerance at the germination period, evaluation of the percentage of reduction in coleoptile length and of coleoptile regrowth is, then, indicated for the identification of cold tolerant genotypes, to be used as genitors. Percentage of reduction in coleoptile length, however, is not viable as a selection criterion in any segregant generation, with the advantage of being a method that involves little space and time for evaluating a high number of plants simultaneously. Recommendation of which generation to select, however, depends on the determination of the inheritance and heritability of the characteristic in cold tolerant genotypes.

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Received January 24, 2003
Accepted November 10, 2003