Growth Response of Hydroponic Rice Seedlings at Elevated Concentrations of Potassium Chloride

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ABSTRACT

A hydroponic experiment was conducted in the greenhouse to observe the effect of potassium chloride (KCl) on growth, macro elements and sodium (Na) concentration of rice seedlings (Oryza sativa L. cv. Hitomebore). The plants were grown up to 14 days after treatments (DAT) treated with 0, 10, 20 and 30 mM KCl. Shoot dry weight (dw) decreased significantly in the 10 mM KCl treatment, indicated that rice seedlings were highly sensitive (glycophytes) to KCl. However, root dw increased in the 10 mM KCl treatment as compared to other treatments, indicated that KCl could enhance root dw at lower concentration. The result indicated that the shoot of rice seedlings was highly sensitive to KCl than that of root. Shoot height, root length and leaf number decreased significantly in the 20 and 30 mM KCl treatments, while the width of leaf blade decreased significantly only in the 30 mM KCl treatment. Considering 10% dw reduction the critical toxicity levels (CTL) of K in shoot was 60.4 mg g⁻¹ dw and 23.3 mg g⁻¹ dw in root. Potassium chloride decreased calcium (Ca), magnesium (Mg) and Na concentrations both in shoot and root, indicated that K had antagonistic relationships with Ca, Mg and Na in rice seedlings. Phosphorus (P) concentration decreased in shoot in the 30 mM KCl treatment indicated that the toxic effete of KCl on P. Potassium concentrations increased in shoot and root with the increasing KCl concentrations in the nutrient solution. The sensitivity response of the elements in rice was Na > Mg > Ca > P under elevated concentrations of KCl.

Key words: Calcium, hydroponic, magnesium, potassium, potassium chloride, rice, sodium.

INTRODUCTION

Some plants could encounter salt stress, but most plants are affected adversely by high levels of salt. Excess salts commonly cause a reduction in growth and yield of most agricultural crops. It was considered that reduction is due to direct effects of ions on photosynthesis that led to reduce growth. The effect of salt on growth reduction is not uniform among ions; some are more toxic than the others. The toxicity of an equamolar concentration of one ionic species is greater than that of a combination of ions (Bernstein, 1975). Although the enzymes of glycophytes (salt sensitive) and halophytes (salt resistant) are equally sensitive to salts, the effects of soil solution salts on plant metabolism vary among ionic species and between glycophytes and halophytes. In glycophytes, Na⁺ salts are more toxic than K⁺ salts, while in halophytes, toxicity follows the general pattern: K₂SO₄ > KCl > Na₂SO₄ = NaCl (Egan and Ungar, 1998).

The mentioned salts could be accumulated in soils and severely inhibit plant growth. They induce plasmolysis (water moves out of the plant into the solution) (Tan, 1998). Intensive research
began in the 1950s, when the U.S. salinity lab showed that some plants can grow faster with low salt content in their tissues and the shoot salts are controlled by the root (Orcutt and Nilsen, 2000). More than 100 plant families contain salt-tolerant species (Aronson, 1989). Most of the families contain less than 10 salt tolerant species, but Chenopodiaceae contains over 350 (Orcutt and Nilsen, 2000). Salts of the soil solution govern the salinity properties of soils. On the basis of salinity tolerance plants can be classified into two broad groups (1) **Glycophytes**: these plants are sensitive to relative low soil salt concentration. Most of the agricultural crops and higher plants are glycophytes. Among crops, maize, onion, citrus, pecan, lettuce and bean are highly sensitive to salt (Taiz and Zeiger, 2002). Glycophytes can be classified as (a) *Euglycophytes* or *true glycophytes* or *strict glycophytes* or *sensitive glycophytes* or *intolertant glycophytes*: a plant whose growth and fitness decreases when expose to any level of salinity (less than approximately 10 mM) (b) *Obligate or Normal glycophytes*: these plants can tolerate a very low salinity but can’t tolerate soil salinity greater than approximately 10 mM (2) **Halophytes**: plants those are able to grow in the presence of high salt condition. These plants often require some salt to grow their best. It is considered that more than one-third of extant plant families have halophytic members and about 250 species of halophytes are found in the world’s salt marshes and desert flats (Glenn, 1997). Halophytes can be classified as (a) *Euhalophytes* or *true halophytes* or *strict halophytes*: halophytes could grow in conditions of extreme high salinity (more than 50 mM) before a reduction in growth or fitness (b) *Obligate halophytes*: plants that require salt in the growth medium to reach their maximum growth potential and have their highest fitness. This type of plant does not attain their highest fitness at low soil salinity (less than 10 mM). Moderately high soil salinity (50 mM) is required to attain maximum fitness (c) *Facultative halophytes*: these plants attain a halophytic nature only after experience moderate salinity in the soils or as an aerosol. Until now, many researches have been done to classify the plants related to salt tolerance activity. Most of which were conducted with Na and its derivatives together with other salt, but the data of KCl research in hydroponic culture were not adequate. Therefore, the present experiment was conducted to define the classification of Hitomebore rice variety together with its macroelements and Na under elevated concentrations of KCl.

**MATERIALS AND METHODS**

**Seedling preparation and plant culture**

Rice seeds were (*Oryza sativa* L. cv. Hitomebore) surface sterilized with 2% (w/v) chlorinated lime ([Ca(OCl)]$_2$) for 45 minutes, rinsed with tap water continuously for 1 hour and soaked between moistened towels covered with wrapping paper at 25°C for 72 hours in an electric incubator. Germinated seeds were placed on plastic net in the seed box containing 2 mM CaCl$_2$ solution in the greenhouse of Iwate University, Morioka, Japan. After 7 days, the chlorinated solution in the seed box was replaced with ½-strength modified Hoagland-Arnon nutrient solution for rice (Hewitt and Smith, 1975). This was done 2 times before starting the treatments. A total of 24 days were required for the seedling preparation. All plants were allowed to grow until 5th full leaf and some plants also contained 6th leaf at emergence stage. After 24 days, seedlings were transplanted in bunches (5 plants were wrapped with sponge rubber to make 1 bunch) and 8 bunches were placed in one pot (capacity 5 liters) filled with full strength rice solution containing 10 µM Fe$^{3+}$-citrate. The plants were treated with KCl up to 14 days after treatments (DAT). The treatments of KCl were 0, 10, 20 and 30 mM. The plants were grown in the greenhouse and the pH of the solutions was 5.5 at the time of solution preparation. The nutrient solutions were renewed every week during the study (July 22, 2005 to August 28, 2005).

**Compositions of full-strength nutrient solution**

The compositions of the full-strength nutrient solutions were 1 mM NH$_4$NO$_3$, 1 mM K$_2$SO$_4$, 0.8 mM MgSO$_4$, 0.5 mM CaCl$_2$, 0.5 mM NaH$_2$PO$_4$, 10 µM MnSO$_4$, 1 µM CuSO$_4$, 1 µM ZnSO$_4$, 3 µM H$_3$BO$_3$, 0.05 µM H$_2$MoO$_4$ and 10 µM Fe$^{3+}$-citrate.

**Sample collection and preparation**

After 14 DAT, 3 bunches were taken from each treatment for determining some growth parameters and P, K, Ca, Mg and Na. The plants were washed with deionized water properly and separated into shoot
and root with sterilized scissors and were dried for 48 hours at 55-60°C in an electric oven (ISUZU, Isuzu Seisakusho Company, Tokyo, Japan). The samples were cut into small pieces and digested with nitric-perchloric acid mixture (Piper, 1942) and analyzed.

**Shoot height, root length and width of leaf blade**

Shoot height was measured from the culm base to the tip of the longest leaf and root length was measured from the root-shoot junction to the tip of the longest root (Abedin and Meharg, 2002). The width of leaf blade was measured in the middle position of fully developed sixth leaf with scale.

**Chlorophyll index (SPAD value)**

The chlorophyll index of fully developed leaves (sixth) on 14 DAT was measured using a SPAD-502 chlorophyll meter (Minolta Camera Company, Tokyo, Japan). In each leaf the chlorophyll index of 5 points were measured and the average was calculated. Means of each bunch were obtained. Average of the data of 3 bunches was calculated.

**Chemical analysis**

Phosphorus was determined colorimetrically using a UV-visible Spectrophotometer (model UV mini 1240, Shimadzu Corp., Kyoto, Japan) at 420 nm wavelength after developing the yellow color with vanadomolybdic acid as described by Barton (1948) and Jackson (1958). Potassium, Ca, Mg and Na were determined with the Hitachi 170-30 atomic absorption spectroscopy (AAS) (Hitachi, Tokyo, Japan).

**Experimental design**

The experiment was a completely randomized block design with 3 replications. Data were analyzed by analysis of variance (SAS, 1990).

**Calculation procedure**

Relative shoot weight in % (RSW) = (shoot weight of a specific treatment/shoot weight of control treatment) × 100. Relative root weight in % (RRW) = (root weight of a specific treatment/root weight of control treatment) × 100. Concentration of elements in plant tissue refers to the element present in one g of sample [mg or µg of the element g⁻¹ dry weight (dw)] while accumulation refers to the specific element present per plant shoot or root (mg or µg of the element plant⁻¹ shoot or root). Uptake is being considered as the total accumulation (accumulation in shoot plus accumulation in root). Translocation (%) could be expressed in nutrient accumulation in shoot × 100/ total accumulation (shoot + root).

**Reagents**

All chemicals used were of analytical reagent grade. All solutions were prepared previously with MQ water [(18.2 MΩ cm⁻¹), purified by Milli-RO 60, Millipore corporation, USA]. Stock solution of 3 M KCl was prepared by dissolving KCl salt (Kanto Chemical Company, Tokyo, Japan) in MQ water. This stock solution was kept at room temperature in an acid washed reagent bottle in the laboratory in normal condition. Working solutions of 0, 10, 20 and 30 mM KCl were prepared every week by diluting the stock solution with deionized water. Glassware used for this experiment was soaked in 0.1 N nitric acid solutions and were rinsed with MQ water.

**RESULTS AND DISCUSSION**

**Visible symptoms**

Severe necrotic symptom (burning of young leaf tips and death associated with discoloration and dehydration of old leaves) was visualized in the leaves of 30 mM KCl treatment. In the 30 mM KCl treatment, the older leaves died completely. Yellowing of younger leaf increased with the severity of KCl concentrations. Greening of the leaves decreased with increasing KCl concentrations in the nutrient solution, this was most probably due to the reduction of chlorophyll content in the leaves. The youngest leaves failed to unfold, leaf size (length and width) decreased drastically and visible growth reductions were found in the 30 mM KCl treatment. Curling of the younger leaves followed by shrivelling occurred. The most common symptom, however, was growth reduction. Potassium chloride toxicity resulted in poor root length, reddish discoloration and felt ‘slippery’ to touch may be
due to deterioration of the middle lamella by excess K. Potassium could interfere physiological availability of Ca (Marschner, 1998) which may lead root function disruption. In this experiment, root length increased in the 10 mM KCl treatment, indicated that low concentration of KCl enhanced root elongation which was most probably due to the positive effect of chlorine (Brady and Weil, 2002). The result of the present study indicated that at high level of availability of KCl resulted in harmful physiological response in the rice seedlings tissues (toxicity range).

Chlorophyll index (SPAD value)

In the present experiment, the chlorophyll index decreased with the increasing KCl concentrations in the nutrient solution (data not shown), indicated that excess KCl reduced the production of chlorophyll content in the leaves.

Dry matter yield

Potassium chloride decreased shoots dw in the 10, 20 and 30 mM KCl treatments. However, root dw increased in the 10 mM KCl treatment, but decreased in the 20 and 30 mM KCl treatments (Fig. 1a). The highest shoot dw was recorded in the control and the lowest was in the 30 mM KCl treatments, respectively. Potassium chloride in the nutrient solution was accountable for 10, 19 and 27% shoot dry matter yield reduction in the 10, 20 and 30 mM KCl treatments, respectively. However, 10% dw increased in root in the 10 mM KCl treatment [this was most probably due to the positive effect of Cl on root length (Brady and Weil, 2002)] while 19 and 24% dw decreased in the 20 and 30 mM KCl treatments, respectively. Our experimental data indicated that the rice seedlings were KCl sensitive (obligate glycophytes) and shoot was more sensitive than that of root. 10–20 mM KCl could be the limiting factor for shoot of rice seedlings in the hydroponic culture. Growth reduction was most probably due to the toxic effects of KCl on enzymatic activity. It is reported that the enzymes of glycophytes and halophytes (Greenway and Osmond, 1972; Greenway and Munns, 1980) are inhibited by salt in vitro equally. We calculated 100, 94, 81 and 73% RSW and 100, 110, 81 and 76% of RRW for 0, 10, 20 and 30 mM KCl treatments, respectively, indicated that with an increase of KCl levels reduced the relative growth.

Shoot height and root length

Shoot height decreased with increasing KCl in the nutrient solution and the lowest value was recorded in the 30 mM KCl treatment. In 10 mM KCl treatment, root length increased significantly as compared to other treatments (Fig. 1b). This increase was most probably due to the positive effect of Cl on root length (Brady and Weil, 2002). However, the root length decreased significantly in 20 and 30 mM KCl treatments. In the higher KCl treatments, Ca concentrations decreased in root (Fig. 2c). The reduction of Ca concentrations might be involved to reduce the root length, because Ca is mainly related to root elongation. Under normal conditions, the cytosol of higher-plant cells contains 100–200 mM K⁺ and 1 to 10 mM Na⁺ an ionic environment in which many enzymes function optimally (Marschner, 1998). In presence of high KCl (30 mM), an abnormally high ratio of K⁺ and high concentrations of total salts may inactivate enzymes and inhibits protein synthesis. At higher concentrations of K⁺, K⁺ may displace Ca²⁺ from the plasma membrane of rice root hairs, resulted in a change in plasma membrane permeability. Cramer et al. (1985) found Ca was replaced by Na in cotton roots and decreased the normal function of root.

Leaf number

Leaf number decreased significantly in 20 and 30 mM KCl treatments. The highest leaf number was recorded in the control treatment and the lowest was in the 30 KCl treatments (Fig. 1c). This was most probably due to the reduced plant growth hormone production or an increase in the production of abscisic acid (ABA) (Nilsen and Orcutt, 1996). This supposition needs to be verified in the future. This reduction of leaf number might be responsible for the reduction of shoot dw (Fig. 1a).

Width of leaf blade

The width of leaf blade decreased significantly in the 30 mM KCl treatment. The highest width of leaf blade was recorded in control treatment and the lowest was recorded in 30 mM KCl.
treatments, respectively (Fig. 1c). This was most probably due to the fact that KCl reduced the activity of plant growth regulators. It was found that protein synthesis is reduced in soybean leaves growing in NaCl saline condition which might be due to chloride toxicity in sensitive species (Marschner, 1998). Therefore, the other possible cause may be reduction of protein synthesis in the higher KCl concentration.

Fig 1. (a) Dry matter yield, (b) shoot height and root length; and (c) leaf number and width of leaf blade of rice seedlings with different levels of KCl. Bars with different letters are significantly different (p<0.05) according to a Ryan-Einot-Gabriel-Welsch multiple range test.
Tiller number

No new tiller was found even in the control treatment during the experiment (38 day after germination) or 14 DAT. Therefore, it was not wise to draw any comment related to effect of KCl on tiller formation. It could only be said that the time duration was not sufficient for the formation of new tiller in this experiment.

Effect of KCl on phosphorus

The concentrations of P did not change in shoot in 10 and 20 mM of KCl treatments as compared to control treatment, but decreased significantly in the 30 mM KCl treatment (Fig. 2a). This reduction was most probably due to severe toxicity of KCl on plant growth. Chlorine is one of the non-metal micronutrients and because of its non-metallic character, it is similar to P. Due to this chemical similarity a competitive process in their uptake could be expected. In the case of root, the concentration decreased significantly in the 10 mM KCl treatment, which was most probably due to dilution effect, because in this treatment root dw and length increased significantly. For other treatments no change of P concentrations were found. Phosphorus concentrations decreased 12% in shoot and 9% in root in 30 mM KCl treatment in the nutrient solution. The accumulation also decreased significantly in shoot and root in the 30 mM KCl treatment which indicated the toxic effect (Table 1). Phosphorus and K ratio in shoot and root decreased significantly with the KCl treatments indicated the negative effect of KCl on P concentration (Table 2). Typical P concentrations in common plants are 2-4 mg g⁻¹ dw (Liao et al., 2004) or 3-4 mg g⁻¹ dw (Mengel and Kirkby, 2001) during the vegetative growth stage, indicated that the P concentrations of the experimental plants was higher (Fig. 2a), this was most probably due to species of rice seedlings. Phosphorus translocation was not much affected in this experiment by the used KCl in the nutrient solution (Table 1).

<table>
<thead>
<tr>
<th>Treatments</th>
<th>P</th>
<th>K</th>
<th>Ca</th>
<th>Mg</th>
<th>Na</th>
</tr>
</thead>
<tbody>
<tr>
<td>(mM KCl)</td>
<td>Shoot</td>
<td>Root</td>
<td>Shoot</td>
<td>Root</td>
<td>Shoot</td>
</tr>
<tr>
<td>0</td>
<td>1.35A</td>
<td>0.26a</td>
<td>4.61C</td>
<td>0.65d</td>
<td>0.44A</td>
</tr>
<tr>
<td>10</td>
<td>1.31A</td>
<td>0.24a</td>
<td>4.74B</td>
<td>0.83b</td>
<td>0.37B</td>
</tr>
<tr>
<td>20</td>
<td>1.06B</td>
<td>0.23a</td>
<td>7.93B</td>
<td>0.93a</td>
<td>0.27C</td>
</tr>
<tr>
<td>30</td>
<td>0.88C</td>
<td>0.18b</td>
<td>8.72A</td>
<td>0.77c</td>
<td>0.25D</td>
</tr>
</tbody>
</table>

Effect of KCl on potassium

Potassium concentrations increased significantly both in shoot and root with the increasing KCl concentrations in the nutrient solution (Fig. 2b). In shoot, the highest concentration was recorded in the 30 mM KCl treatment. Not only concentration but also accumulation increased in shoot with treatments (Table 1). In the highest concentration of KCl in the nutrient solution, the translocation of K from root to shoot did not increase markedly (Table 1). The increase of K concentration and accumulation in shoot and in root might be due to “luxury consumption” of K (Marschner, 1998) which is a normal phenomenon in high level of K supply. The K content increased in various organs [except grains and seeds which maintain relatively constant K content of 0.3% of the dw] by increasing the K supply to plant root (Marschner, 1998). The value of K concentration is 1-4% (10-40 mg K g⁻¹ dw) of dw in healthy leaf tissue of most plants (Brady and Weil, 2002). However, it was also pointed out that K requirement for optimal plant growth is in the range of 2-5% (20-50 mg g⁻¹ dw).
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dw) of plant vegetative parts, fleshy fruits and tubers (Marschner, 1998). In the highest concentration of KCl, the plants absorb high concentrations of K, resulted in the toxicity symptoms. At high level of availability of an element, the plant will take up a large amount of the nutrient elements for its own good (toxicity range), result harmful physiological reactions in the plant cell. In this experiment, the K concentration in shoot of control seedlings was 35.9 mg g\(^{-1}\) dw, indicated the normal concentration of K in shoot tissues. It was also reported that K concentration in shoot of hydroponic rice was 44.1 mg g\(^{-1}\) dw (Shaibur et al., 2006) and 60.2 mg g\(^{-1}\) dw in hydroponic barley (Alam et al., 2001) of control plants.

**Effect of KCl on calcium**

Calcium concentrations decreased both in shoots and roots with the increasing KCl concentrations in the nutrient solution. The significant reductions were obtained in 20 and 30 mM KCl treatments (Fig. 2c). It is well known that KCl in solution is dissociated as K\(^+\) and Cl\(^-\) ions. As K\(^+\) and Ca\(^{2+}\) both are cations, therefore, antagonistic relationships between them could be possible. Our result showed that the antagonistic relationship between K\(^+\) and Ca\(^{2+}\); and the Ca concentrations decreased by 22% in shoot and 27% in root in the 30 mM KCl treatment. When the K supply is abundant in the rooting medium "luxury consumption" of K often occurs which deserve attention for its possible interference with the uptake and physiological availability of Ca and Mg (Marschner, 1998). Calcium stabilizes cell membranes by bridging phosphate and carboxylate groups of phospholipids (Caldwell and Haug, 1981) and preferentially protein at membrane surfaces (Legge et al., 1982). There can be an exchange between Ca at these binding sites and other cations (eg. K\(^+\), Na\(^+\) or H\(^+\)) resulted in Ca concentrations decreased. Calcium accumulations were also decreased significantly by the KCl treatments but the translocation was not affected much (Table 1). Calcium concentration in plant tissues varies between 0.1->5% (1-50 mg Ca g\(^{-1}\) dw) of dw depending on the growing conditions, plant species and plant organ (Marschner, 1998). The experimental plants contained 3.44, 3.12, 2.64 and 2.68 mg Ca g\(^{-1}\) dw in shoot tissues in 0, 10, 20 and 30 mM KCl treatments, respectively, indicated that the normal levels of Ca concentrations. The highest Ca/K ratio was obtained (both in shoot and root) in control treatment and the lowest was in 30 mM KCl treatment, indicated that the lowest Ca concentration was in the highest KCl concentration in the nutrient solution. The hydrated K\(^+\) has a radius of 0.331 nm, but Ca\(^{2+}\) has the ionic radius of 0.099 nm (Marschner, 1998). Therefore, K strongly substitutes or competes with Ca\(^{2+}\) in various reactions involving either of these ions.

**Effect of KCl on magnesium**

Marginal concomitant relationship was found between K and Mg in shoot in the 10 mM KCl treatment (Fig. 2d). It may be due to the concentration effect because shoot dw decreased significantly in this treatment (Fig. 1a). Magnesium concentrations decreased significantly in shoot in 20 and 30 mM treatments and in root in 10, 20 and 30 mM KCl treatments. This is logical because K\(^+\) and Mg\(^{2+}\) both are cations, therefore, antagonistic relationship between them might be occurred. Some reports indicated that the Mg uptake can strongly be depressed by K\(^+\) and NH\(^4\) (Kurvits and Kirkby, 1980), Ca\(^{2+}\) and Mn\(^{2+}\) (Heenan and Campbell, 1981) as well as by H\(^+\). Magnesium concentration decreased by 30% in shoot and 33% in root in 30 mM KCl treatment in the nutrient solution. The data presented in Table 1 showed that the accumulation of Mg was also depressed by the used KCl treatments, indicated that the antagonistic relationship between K and Mg, however, the translocation of Mg was not affected by KCl (Table 1). The Mg and K ratio in shoot and in root decreased significantly with the KCl treatments (Table 2). In presence of excess available K, K may be taken up as high level (Marschner, 1998) which interfering the uptake and physiological availability of Mg, resulted in the decrease in the Mg/K ratio in the present experiment. In leaf tissues, the threshold value for the occurrence of Mg-deficiency symptoms is in the region of about 2 mg g\(^{-1}\) dw (Mengel and Kirkby, 2001). The test plants contained 3.18, 3.40, 2.54 and 2.22 mg Mg g\(^{-1}\) dw in 0, 10, 20 and 30 mM KCl treatments, respectively, indicated that the value in shoot tissues was over the critical deficiency level of Mg. It was also reported that the Mg requirement for optimum plant growth is in the range of 0.15-0.35% (1.5 -3.5 mg Mg g\(^{-1}\) dw) of the vegetative parts (Marschner, 1998), indicated that the shoot of the experimental plants contained more or less normal concentrations of Mg. Potassium has a hydrated radius of 0.331 nm and Mg\(^{2+}\)
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has the radius of 0.065 nm (Marschner, 1998). As a result, Mg could be replaced strongly by K in various reactions in this present experiment.

Fig 2. Effect of KCl on the concentration of (a) P, (b) K, (c) Ca and (d) Mg in shoot and root of rice seedlings. Bars with different letters are significantly different (p<0.05) according to a Ryan-Einot-Gabriel-Welsch multiple range test.
Table 2. Concentration ratio of elements in shoot and in root of rice seedlings grown in nutrient solution at different levels of KCl

<table>
<thead>
<tr>
<th>Treatments (mM KCl)</th>
<th>P/K Shoot</th>
<th>P/K Root</th>
<th>Ca/K Shoot</th>
<th>Ca/K Root</th>
<th>Mg/K Shoot</th>
<th>Mg/K Root</th>
<th>Na/K Shoot</th>
<th>Na/K Root</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.293 A</td>
<td>0.401 a</td>
<td>0.096 A</td>
<td>0.023 a</td>
<td>0.087 A</td>
<td>0.042 a</td>
<td>0.065 A</td>
<td>0.279 a</td>
</tr>
<tr>
<td>10</td>
<td>0.181 B</td>
<td>0.291 b</td>
<td>0.054 B</td>
<td>0.015 b</td>
<td>0.054 B</td>
<td>0.031 b</td>
<td>0.027 B</td>
<td>0.114 b</td>
</tr>
<tr>
<td>20</td>
<td>0.134 C</td>
<td>0.241 c</td>
<td>0.034 C</td>
<td>0.013 b</td>
<td>0.033 C</td>
<td>0.021 c</td>
<td>0.016 C</td>
<td>0.106 b</td>
</tr>
<tr>
<td>30</td>
<td>0.101 D</td>
<td>0.232 c</td>
<td>0.029 D</td>
<td>0.011 c</td>
<td>0.024 D</td>
<td>0.018 d</td>
<td>0.015 C</td>
<td>0.078 c</td>
</tr>
</tbody>
</table>

*Means followed by the different letters in each column are significantly different (p = 0.05)*

Effect of KCl on sodium

Sodium concentrations decreased significantly in shoot and root with the increasing KCl concentrations in the nutrient solution (Fig. 3), indicated that the severe antagonistic relationship between these two elements. Higher plants have developed high selectivity in the uptake of K as compared with that of Na and this is particularly obvious in transport to shoot (Marschner, 1998). For this reason “luxurious consumption of K” was taken place in the high KCl concentrations in the nutrient solution. Potassium has a more or less similar hydrous radius (0.331 nm) like Na⁺ (0.358 nm) (Marschner, 1998). In another way, it has also been pointed out that Na has the ionic radii of 1.65~2.05 Å while in K the ionic radii is 2.350~2.66 Å (Biggin et al., 2001). Therefore, strong competition between them could be possible, resulted in lower concentrations of Na in shoot and root in this experiment (Fig. 3). Moreover, both are positively charged cations. In some natrophilic plants, K can be replaced when Na content is high (Marschner, 1998). Similarly, opposite result could be expected in higher KCl solution in this present experiment. In sugar beet Na is readily translocated to the shoot where it replaces most of the K (Marschner, 1998). In this experiment, not only concentrations but also accumulations and translocations of Na decreased with the increasing KCl concentrations in the nutrient solution (Table 1). Sodium concentration decreased by 41% in shoot and 56% in root in the 30 mM KCl treatment. Potassium concentrations increased by 75%, 114% and 159% in shoot but the values were 13, 72 and 56% in root in 10, 20 and 30 mM KCl treatments, respectively. The highest Na and K ratio in shoot and in root were obtained in the control treatment (Table 2). The ratio decreased gradually and significantly and the lowest value was in the 30 mM KCl treatment. This result indicated that the highly antagonistic relationships between K and Na. When salinity is caused by Na salt, it may decrease K content in plant tissue by competition, as both of them have similar ionic radii at hydrated condition.

![Graph](image-url)

**Fig 3.** Effect of KCl on the concentration of Na in shoot and root of rice seedlings. Bars with different letters are significantly different (p<0.05) according to a Ryan-Einot-Gabriel-Welsch multiple range test.
Critical toxicity levels (CTL) of K in shoots and roots

Considering 10% dw reduction the CTL of K was calculated as 60.4 mg g\(^{-1}\) dw in shoot and 23.3 mg g\(^{-1}\) dw in root, though the CTL were suggested to be very much dependent on the species and physical conditions of plants, varieties, environmental conditions especially temperature, nutrient compositions and methodology of the experiments. The CTL were calculated from the polynomial two order growth curves (Fig. 4ab) where the equations were \(y = -0.0071x^2 - 2.1289x + 733.86\) and \(y = -0.1972x^2 + 2.9083x + 224.02\) for shoot and root, respectively. For shoot, the highest growth was in control treatment and the value was 643.6 mg/5 plants and the 10% dw reduction was calculated from this value. In case of root, the value of control treatment was 205.4 mg/5 plants and the value was used for the determination of CTL in root. By using the similar equations the CTL of arsenic (As) were reported by the present author in rice, sorghum and barley (Shaibur et al., 2008).

![Fig 4. Two order polynomial growth curve (a) shoot and (b) root of rice seedlings with different KCl concentrations in plant tissues](image)

CONCLUSION

The dry weight decreased significantly in shoot in the 10 mM KCl treatment, but in root the significantly reductions were obtained in 20 and 30 mM KCl treatments. Shoot height, root length and leaf number decreased significantly in 20 and 30 mM KCl treatments, but width of leaf blade decreased in the 30 mM KCl treatment. Therefore, rice plants could usually be classified as obligate glycophytes where shoots were more sensitive to KCl than that of roots. The critical toxicity levels (CTL) of K were 60.4 mg g\(^{-1}\) dw in shoot and 23.3 mg g\(^{-1}\) dw in root. From this experiment it could be concluded that K\(^+\) shows the antagonistic relationships with Ca, Mg and Na in rice seedlings. The decrease of P concentration was most probably due to the toxic effect of 30 mM KCl treatment on rice seedlings. Phosphorus, Ca, Mg and Na concentrations decreased in shoot by 12, 22, 30 and 41% in the 30 mM KCl treatment, but for root the values were 9, 27, 33 and 56%, respectively. Considering the reduction value of concentrations, the sensitivity of these elements can be arranged as Na> Mg>Ca>P both for shoot and root in rice seedlings when KCl is present at 30 mM level. From our result it was suggested that small amount of KCl (10 mM) could be the good fertilizer for the rooting crops as the root growth increased significantly in the 10 mM KCl treatment, though this suggestion needs to be verified. In most cases it was found that KCl decreased the concentrations of P, Ca and Mg. The concentrations of macroelements were within the normal concentrations of plant nutrition but the plant growth decreased. Therefore, it could be suggested that the reduction of growth was associated with the reduction of macroelements concentrations together with the production of toxic biochemical substances which needs to be investigated.

LITERATURE CITED


