Effect of pre-harvest Potassium Treatment on Stem-end Rot Disease Development of Mango (*Mangifera indica* L.)cv. TomEJC during Fruit Ripening

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Abstract

Mango (*Mangifera indica* L.) cv. TomEJC has utmost potential in export market due to its unique quality. Post-harvest disease of mango such as stem-end rot can cause considerable losses of fruits and could therefore be considered as a great threat to local and export market. This disease is controlled by application of fungicides from the time of flowering in cultivations where this disease is severe. However, the use of fungicides could lead to hazardous effects such as oncogenic risks on the consumers. Therefore, search for alternative measures for the management of stem-end rot disease is essential. In this study, an attempt was made to find out the effect of pre-harvest application of KCl on controlling stem-end rot disease development of ripe fruit. KCl at concentrations of 1 gl⁻¹, 2 gl⁻¹ and 4 gl⁻¹ were sprayed on fruit when fruits were immature. All fruits were bagged with paper bags and fruits were harvested at the correct maturity stage. Then all fruits were inoculated with Botryodiplodia theobromae mycelial plug and allowed for natural ripening. A significant difference was observed in fruit length, fruit width and fruit volume in KCl- treated fruits when compared to controls. Highest fruit volume was observed in 1 gl⁻¹ KCl- treated fruits and the lowest amount of total soluble solids was observed in 2 gl⁻¹ KCl- treated fruits. An increase in lightness and yellowness of peel color in KCl- treated fruits when compared to control II (non-inoculated, non-treated) was evident, indicated by increasing L*, b* and chroma values. However, no significant difference in hue values of peel color was observed among treatments. Flesh color showed a different response to treatments and no difference in chroma value was observed among treatments. The highest L* value and hue value was observed in 2 gl⁻¹ KCl- treated fruits. Incidence and severity of stem-end rot was significantly decreased by 2 gl⁻¹ KCl treatment without drastically affecting the physico-chemical properties of fruits. Vast research has shown that potassium and chloride fertility have been effective in reducing crop injury from diseases. Different mechanisms such as nutritional effects, changes of the
host-pathogen environment and production of disease inhibitory compounds could be some possible ways of action of both K and Cl in modifying disease resistance or susceptibility. Since 2 gl$^{-1}$ KCl was able to suppress stem-end rot disease development in inoculated fruits, this concentration can be recommended as a field application to control this disease in mango variety tested after repeating the same experiment at different locations.

**Keywords:** Cultivar TomEJC, Botryodiplodia theobromae, post-harvest loss, stem-end rot disease, disease severity, disease incidence, potassium chloride

**Introduction**

TomEJC Mango (*Mangifera indica* L.) is the world’s biggest, juiciest mango. It is a beautiful golden orange fruit with an unblemished skin. It has an excellent flavor, low fibre content and smooth flesh. The fruit ripens slowly giving time for the produce to reach the market (Wijesinghe *et al*., 2011). TomEJC is well adapted to the dry zone in Sri Lanka. The existence of this cultivar could be identified as a new variety having distinct properties when compared to the other standard recommended Sri Lankan varieties.

Domestic and international trade of fresh mango has been limited by its perishable nature and its susceptibility to post-harvest diseases. Post-harvest diseases of mango reduce fruit quality and cause severe losses because they leave them completely unmarketable (Bally *et al*., 2009; Barkai-Golan, 2001; Narayanasam, 2006). Major post-harvest diseases of mango that deteriorate the fruit quality include anthracnose and stem-end rot disease (Jefferies *et al*., 1990; Crane and Campbell, 1991).

Next to anthracnose, stem-end rot is considered a major problem limiting the storage and shelf life of mango fruits. Stem-end rot is a disease of importance in harvested fruit which reduces consumer acceptability. A number of fungi including *Dothiorella dominicana*, *Phomopsis* spp., *Botryodiplodia theobromae* and *Lasiodiplodia theobromae* cause stem end rot in mango ([https://www.daf.qld.gov.au/plants/fruit-and.../a-z-list.../stem-end-rot](https://www.daf.qld.gov.au/plants/fruit-and.../a-z-list.../stem-end-rot)). Disease symptoms develop around the stem end of the fruit as it begins to ripen after harvesting. Fungal spores are prevalent on dead leaves and twigs in orchards. As the spores spread on to flowers and fruit before harvesting, the fungi may form latent infection on fruit. Symptoms become clear as the fruit ripens. At the stem end of the fruit, brownish patches begins to develop. Due to this, both the peel and the inside flesh begins to rot. Infected fruit emanates an undesirable odour and fruit becomes unmarketable.

Post- harvest losses of fruits and vegetables that occur due to diseases in developed countries range from 5-25%, while in developing countries it is 20-50% depending upon the commodity (Kader, 1992). However, post-harvest losses of mango accounts for around 41% of the total produce in Sri Lanka (Karunanayaka & Adikaram, 2011).
Control of post-harvest diseases of mango fruit mostly depends on the combination of pre and post-harvest fungicide treatment. However, increasing consumer attention on health risks and environmental hazards associated with the use of pesticides (Wilson et al., 1994), fruits and vegetables free from pesticide residues, the legislative restriction aiming at increasing food security, the development of pathogen strains resistant to few admitted post-harvest fungicides and the need for high quality products have increased the search for control measures alternative to chemical fungicides (Sanzani et al., 2009).

Recently, disease control by inducing host resistance and activating the defense mechanisms in plants (especially herbaceous plants) and harvested fresh produce (Johnson & Hofman, 2009) have been reported.

Many species of potassium-deficient plants are susceptible to frost damage and certain diseases than plants with adequate potassium levels. Increased disease resistance associated with adequate potassium levels indicate that potassium has a role in providing disease resistance and increasing the potassium levels of deficient plants have been shown to decrease the intensity of many diseases (http://en.wikipedia.org/wiki/Potassium_deficiency_%28plants%29).

The role of K in crop resistance to diseases was extensively reviewed by Perrenoud, (1990) and an inverse relationship was found between available soil K and the severity of disease caused by bacteria and fungi. In potatoes, K fertilization was found to decrease the incidence on several diseases, such as late blight (Phytophthora infestans), dry rot (Fusarium spp.), powdery scab (Spongospora subterranea) and early blight (Alternaria solani) (Perrenoud, 1990; Marschner, 1995). Potassium applications resulted in suppression of diseases such as Tikka leaf spot (Cercospora archidicola Hori.) in groundnut (Umar et al., 1997) and leaf spot disease in cotton (small brown lesions caused by Cercospora, Alternaria and Stemphylium) (Harris, 1997). Potassium fertilizers such as NPK (nitrogen-phosphorus-potassium fertilizers) applied as foliar sprays were highly effective inducers of systemic protection against powdery mildew in cucumber, mango, nectarines and grapes (Reuveni and Reuveni, 1995 a & 1995 b). Further, high K levels of fertilizer were shown to reduce severity of anthracnose of banana and mango in Sri Lanka (Coates, 2002-2007). High doses (three times the recommended level by Department of Agriculture) of potassium supplied as a fertilizer to soil (muriate of potash) effectively controlled stem-end rot in karutha colombon mango (Karunanayake, 2008).

In this study, an attempt was made to find out the possibility of using KCl as an effective agent of controlling stem-end rot disease of “TomEJC” mango.
Materials and Methods

Mango cultivar “TomEJC” was used for this study and the experimental plot was situated at Ellawala farm, Galkiriyagama, Dambulla. Orchard management was done using standard agronomic practices recommended by Department of Agriculture, Sri Lanka (http://www.doa.gov.lk/index.php/en/crop-recommendations/1087).

Three different concentrations of potassium chloride (KCl) were sprayed on mango fruits when they were at egg size (one month after fruit set in the tree). The KCl concentrations used were 1 gl⁻¹, 2 gl⁻¹ and 4 gl⁻¹. Each concentration was sprayed once to ten mango fruits until the entire fruit was wet and the solution drained off from the fruit (approximately 10 ml) in three different trees. Twenty fruits on another tree which was sprayed with sterile distilled water without KCl served as controls (control I- non-treated, inoculated and control II-non-treated, non-inoculated). After spraying, all the treated and control fruits were labeled and covered using special paper bags (out side-brown color, in side-black) imported from Thailand (Sarananda, unpublished data). Fruits were harvested three months after bagging and brought into the laboratory at The Open University of Sri Lanka and experiments were carried out to examine the physico-chemical properties and disease incidence/disease severity. Peel and flesh color measurements were done at the Institute of Post-harvest Technology, Anuradhapura and the determination of potassium content in the fruit peel was carried out at Horticultural Crops Research and Development Institute, Gannoruwa.

Disease incidence and severity of stem-end rot disease

Disease incidence and severity was assessed by artificial inoculation of mango fruits with Botryodiplodia theobromae. The mango fruits were surface sterilized using 70% v/v ethanol and washed with sterilized distilled water and air-dried in a lamina flow. Then stalks were removed and stem-ends were gently injured with a sterile scalpel. Then the injured stem-end of treated and control fruits was inoculated (plugged) with 4 mm diameter mycelial discs of pure culture of Botryodiplodia theobromae. Thereafter, inoculated fruits were arranged randomly in a humid chamber (28 °C and 100% RH). Disease incidence was recorded daily and disease severity was assessed by measuring the lesion area (mm²) of infected fruits using a mm graph paper from day 5 to day 9 after inoculation (Karunanayaka and Adikaram, 2011).

Physico-chemical properties

Physico-chemical properties such as peel/flesh color, fruit volume, fruit size and total soluble solids of non-treated and treated fruits were measured after inoculation with Botryodiplodia theobromae.
Fruit size

Fruit length (from stalk end to the apex) and width (at the widest position) were measured using a Vernier caliper (Karemera and Habimana, 2014).

Fruit volume

Fruit volume was measured using water displacement method. Fruits were dipped in a water beaker and displaced water was collected and the volume was measured using a measuring cylinder. This volume was taken as the fruit volume (Saranada, unpublished data).

Total soluble solids (TSS)

A few drops of the filtrate of diluted juice obtained from treated and control fruits were used to measure the TSS using a hand-held refractometer (model-HR-5 A022-1, Kyowa optical Co Ltd., Tokyo, Japan). The reading obtained was multiplied by a dilution factor (DF) to calculate the actual TSS content of the pulp (DF= weight of sample + volume of water added/weight of sample), expressed as Brix (Sarananda & Wijerathnam, 1994).

Peel color and flesh color

Peel color and flesh color were measured objectively using a Minolta Chromameter (Model CR-400, Minolta camera Co. Ltd., Osaka, Japan.). Physical changes in relation to peel color and flesh color were recorded in numerical notation system as L*, a* and b* where L* indicates lightness or darkness (black-0 and white -100), a* ranges from negative values for green to positive values for red and b* ranges from negative values for blue to positive values for yellow. L*, a* and b* values were converted to hue (h°) value and Chroma (C) (McGuire, 1992). The Chroma value \{(C) = (a*^2 + b*^2)^{1/2}\} indicates the strength of color. The hue value was measured as the hue angle (h° = arctangent (b*/ a*) of target color). The hue angle was expressed in degrees: 0° = (red), 90° = (Yellow), 180° = (green), 270° = (blue) (Jha, 2010).

The potassium content in fruit peel

The residual potassium content in fruit peel was measured using the dry ashing procedure. The concentration of potassium in the diluted solution was determined by the flame emission spectrophotometer using the method described by Senevirathna & Daundasekera (2010).

Peels were obtained separately from three randomly selected fruits of each treatment and controls. Each peel sample was air- dried and ground into a fine powder in a domestic grinder. Subsequently, 0.5 g of each ground sample was
ashed in a muffle furnace (Lenton thermal designs, England) at 500 °C for 4 to 5 hours until the ash turned white color. The powdery white dry ash residue obtained was wetted with 2.5 ml concentrated nitric acid and volume was brought up to 25 ml by adding de-ionized distilled water in a volumetric flask. The suspension was swirled for two minutes and filtered through Whatman No. 01 (5.5 cm diameter) filter paper to obtain digested (wet) samples for potassium analysis. A dilution series of the sample solutions were prepared using the above 25 ml sample solution. One (1.0) ml of each sample solution was incorporated into volumetric flasks and the volume was brought up to 50 ml by adding de-ionized distilled water. The residual potassium content of the peel was measured using the flame emission spectrophotometer (Corning, CR 410) and was expressed as percentage potassium.

**Research design and statistical analysis**

Ten replicate fruits were used to assess disease incidence and disease severity, color measurement of flesh and peel, physico-chemical properties and the residual potassium content of mango peel. Data were analyzed using one-way analysis of variance (ANOVA) using SAS V. 9.2 (SAS, 2008), SPSS 16.0 (SPSS Inc., Chicago, IL, USA) to test the difference among treatments, considering all variables simultaneously and Tukey’s Honest Significant Difference (HSD) Test to evaluate the level of effects for all pair-wise comparisons. Canonical Discriminate Analysis was used to represent the linear relationship of the colorimeter variables.

**Results**

**Effect of potassium on disease incidence and severity of stem-end rot disease of ‘TomE JC’ mango**

Disease incidence was reduced to 80% in 2 gl⁻¹ KCl-treated fruits and 40% in 4 gl⁻¹ KCl-treated fruits (Figure 01). 1 gl⁻¹ KCl was less effective than the rest of the concentrations and disease incidence was same in 1gl⁻¹ KCl-treated fruits and the control fruits (control I).

Disease severity in 1 gl⁻¹ KCl-treated fruits was higher than control fruits (control I and II) (Table 01). The concentration of 2 gl⁻¹ KCl-treated fruits showed the lowest disease severity until day 9 from the inoculation (Table 01) and in 2 gl⁻¹ KCl-treated fruits, the lesion development was delayed until day 7. Therefore, 2 gl⁻¹ KCl treatment is the most successful concentration to control stem-end rot disease severity than other concentrations (Table 01).
Table 01: Mean lesion area (mm²) of Stem-end rot disease (Botryodiplodia theobromae) of mango cv. TOMEJC.

| Trt          | Day 05     | Day 06     | Day 07     | Day 08     | Day 09     |
|--------------|------------|------------|------------|------------|------------|
| Control I    | 18.60±(3.79)| 43.20±(10.36)| 62.40±(16.03)| 77.40±(18.78)| 117.40±(29.14)|
| Control II   | 10.20±(4.72)| 13.60±(6.91)| 70.00±(55.77)| 24.40±(13.85)| 29.20±(16.59)|
| 1 gl⁻¹       | 39.20±(21.63)| 52.20±(25.09)| 82.80±(22.47)| 108.80±(25.19)| 138.20±(13.28)|
| 2 gl⁻¹       | (0.00)±(0)  | (0.00)±(0)  | 10.60±(10.60)| 13.00±(13.00)| 13.20±(13.20)|
| 4 gl⁻¹       | 6.00±(2.45) | 8.00±(3.39) | 10.20±(4.41)| 22.00±(9.17)| 25.00±(10.72)|

Mean (n = 10) ± Standard error. Values with the same letters in the same column are not significantly different at (p ≥ 0.05) by Tukey’s Honest Significant Difference (HSD) Test. Control I = inoculated, non-treated, Control II = non-inoculated, non-treated. Trt = Treatment/control.

Effect of potassium on physico-chemical properties of TomEJC’ mango

Fruit length and fruit width slightly changed with increase in KCl concentration (Table 02). The highest fruit volume was observed in concentration of 1 gl⁻¹ KCl-treated fruits and the lowest total soluble solids (TSS) was observed in 2 gl⁻¹ KCl treated fruits (Table 02). The percentage of residual potassium increased with increase in KCl concentration. The maximum level of K was observed in fruits treated with 4 gl⁻¹ KCl which was (0.61± 0.00). The percentage of residual potassium observed in control II and 2 gl⁻¹ KCl-treated fruits was equal (Table 02).
It was observed that L*, b* and chroma values for peel color of KCl- treated fruits showed an increase when compared to control II (non-inoculated, non-treated) indicating a slight increase in lightness and yellowness (Table 03). However, hue values for treatments and controls ranged between 86.53 to 77.17 and was not statistically significant (Table 03).

Flesh color showed a different response to treatments and no difference in chroma value was observed among treatments (Table 03). The highest L* value and hue value for flesh color were observed in 2 gl−1 KCl- treated fruits when compared to control I. Increase in b* value for flesh color in 1 gl−1 and 2 gl−1 KCl- treated fruits when compared to control I indicated a slight increase in yellowness in flesh.

Table 02: Changes in physico-chemical properties of mango cv. TomEJC.

| Treatment | fruit length | fruit width | Fruit volume | TSS | K (%) |
|-----------|--------------|-------------|--------------|-----|-------|
| Control I | 13.04±(0.22) | 8.92±(0.15) | 526.00±(0)   | 14.00±(0) | 0.41±(0) |
| Control II| 15.50±(0.23) | 9.88±(0.12) | 554.80±(11.36) | 14.00±(0) | 0.54±(0) |
| 1gl−1     | 14.62±(0.37) | 9.88±(0.12) | 656.00±(23.79) | 14.00±(0) | 0.47±(0) |
| 2gl−1     | 14.64±(0.29) | 9.82±(0.15) | 625.00±(11.18) | 11.50±(0) | 0.54±(0) |
| 4gl−1     | 13.78±(0.58) | 9.34±(0.28) | 489.00±(30.10) | 14.00±(0) | 0.61±(0) |

Mean (n = 10) ±Standard error. Values with the same letters in the same column are not significantly different at (p ≥ 0.05) by Tukey’s Honest Significant Difference (HSD) Test. Control I = inoculated, non-treated, Control II = non-inoculated, non-treated.

Table 03: Changes in L*, a*, b* values of peel and flesh color of mango cv. TomEJC.

| Treatment | Peel Color | L*          | a*          | b*          | Chroma      | Hue         |
|-----------|------------|-------------|-------------|-------------|-------------|-------------|
| Control I | 59.41±(2.09)| 5.24±(1.41) | 48.57±(2.51)| 26.90±(1.18)| 83.63±(1.91)|
| Control II| 51.25±(3.02)| 7.33±(0.41) | 34.82±(4.38)| 21.07±(2.19)| 77.17±(2.30)|
| 1gl−1     | 61.87±(1.09)| 6.48±(2.25) | 49.83±(2.54)| 28.16±(2.09)| 82.84±(2.26)|
| 2gl−1     | 58.28±(2.18)| 2.53±(0.22) | 42.68±(2.89)| 22.60±(1.43)| 86.53±(0.46)|
| 4gl−1     | 61.52±(1.03)| 5.28±(0.45) | 48.78±(1.47)| 27.03±(0.83)| 83.82±(0.47)|

| Treatment | Flesh Color | L*          | a*          | b*          | Chroma      | Hue         |
|-----------|-------------|-------------|-------------|-------------|-------------|-------------|
| Control I | 66.86±(1.72)| 9.06±(0.71) | 60.69±(1.22)| 34.88±(0.28)| 81.45±(0.83)|
| Control II| 68.72±(1.48)| 8.80±(0.99) | 65.79±(1.13)| 37.29±(0.91)| 82.41±(0.79)|
| 1gl−1     | 69.70±(2.25)| 7.63±(3.29) | 68.07±(1.79)| 37.85±(2.34)| 83.83±(2.57)|
| 2gl−1     | 70.42±(1.31)| 3.88±(1.62) | 65.34±(1.26)| 34.61±(1.38)| 86.69±(1.35)|
| 4gl−1     | 64.21±(2.71)| 9.60±(1.37) | 59.44±(0.93)| 34.52±(0.69)| 80.81±(1.36)|

Mean (n = 10) ±Standard error. Values with the same letters in the same column are not significantly different at (p ≥ 0.05) by Tukey’s Honest Significant Difference (HSD) Test. Control I = inoculated, non-treated, Control II = non-inoculated, non-treated.
Discussion

A considerable reduction in disease incidence was observed in 2 gl\textsuperscript{-1} KCl-treated fruits. The highest reduction in disease severity was also observed in 2 gl\textsuperscript{-1} KCl-treated fruits, and the lesion development was delayed until day 7. However, 1 gl\textsuperscript{-1} KCl was not effective in reducing disease incidence and disease severity when compared to the rest of the concentrations tested.

K has been widely used to control many diseases. Vast research has shown that incidence and rate of development of diseases may be reduced by an adequate and balanced mineral nutrition in many crops. In particular, K and Cl fertility have been effective in reducing crop injury from diseases (Magen and Imas, 2004). Potassium fertilizers such as (NPK) applied as foliar sprays were highly effective inducers of systemic protection against powdery mildew in cucumber, mango, nectarines and grapes (Reuveni and Reuveni, 1995 a & 1995 b) and high doses (three times the recommended level by Department of Agriculture) of potassium supplied as a fertilizer to soil (muriate of potash) effectively controlled stem-end rot in Karutha Colombon mango (Karunanayake, 2008).

According to Cooke et al. (1997), pre-harvest trunk injection of potassium phosphonate when fruits were hen-egg-size and injection of 400 g potassium phosphonate/litre (15 ml/m canopy diameter) after storage at 22 °C for 20 days significantly reduced post-harvest stem-end rot (caused by Dothiorella dominicana and Phomopsis mangifera) in fruits of cultivar Kensington Pride.

The mechanisms involved with increased host resistance and potassium include a decreased cell permeability and decreased susceptibility to tissue penetration. Silica, which is accumulated in greater quantities when adequate potassium is present, is incorporated into cell walls, strengthening the epidermal layer which functions as a physical barrier to pathogens. Potassium has also been implicated to have a role in the proper thickening of cell walls (Datnoff et al., 2007).

It has been found that for many species, potassium-deficient plants are more susceptible to frost damage and certain diseases than plants with adequate potassium levels. Increased disease resistance associated with adequate potassium levels indicates that potassium has a role in providing disease resistance, and increasing the potassium levels of deficient plants have been shown to decrease the intensity of many diseases. In agriculture, some cultivars are more efficient at K uptake due to genetic variations and often these plants have increased disease resistance.(Datnoff et al., 2007).

Potassium deficiency has been found to be linked to diseases in a number of temperate crops (Palti, 1981) and a high K supply can improve resistance of plants to fungal and bacterial pathogens (Marschner, 1995; Perrenoud, 1977;
The mechanism of resistance in some disease-resistant genotypes might be related to a greater efficiency in K uptake (Prabhu et al., 2007).

K-deficient plants have impaired protein synthesis and accumulate simple N compounds such as amides which are used by invading plant pathogens (Marschner 1995). K-deficiency increases the concentration of soluble sugars in leaf tissues providing a substrate for many pathogens (Perrenoud, 1990).

Further, N/K ratio may also affect disease resistance. When this ratio is too high, cells have displayed thinner cell walls and weaker membranes which are more prone to pathogen attack (Perrenoud, 1990). For similar reasons, cereals may become more prone to lodging. A low potassium/chloride (K/Cl) ratio in plant tissues, which might result from the application of chloride-containing compounds such as ammonium fertilizers, may predispose plants to disease e.g. wheat rust caused by *Puccinia* spp. or other diseases (Prabhu et al., 2007; Jones et al., 1989).

It is likely that the susceptibility of tropical perennial crops to some pathogens is also increased under conditions of K deficiency. In a study on tea plants, for example, a high K supply reduced nematode and borer damage (Muraleedharan and Chen, 1997). Another study reported that supplying K reduced *Fusarium* wilt in oil palm (Turner et al., 1970). However, few research studies have been conducted that could confirm a link between K nutrition and disease incidence or severity in tropical perennial crops.

Present study indicates that the pre-harvest application of KCl is effective in controlling stem-end rot disease of “TomEJC” mango. KCl concentration of 2 gl⁻¹ was the most effective in reducing the stem-end rot disease incidence and severity. The reduction in disease incidence and severity may be due to the production of anti-fungal compounds which in turn increase the disease resistance of cultivar ‘TomEJC’ and delay the appearance of post-harvest symptoms.

Further, the fruits treated with 2 gl⁻¹ KCl possess residual K which is similar to the non-treated fruits (control II) and the lowest amount of total soluble sugar content.

b* values for peel color of KCl-treated fruits showed an increase when compared to control II (non-inoculated, non-treated) indicating a slight increase in yellowness. Increase in b* value for flesh color in 1 gl⁻¹ and 2 gl⁻¹ KCl-treated fruits when compared to control I also indicated a slight increase in yellowness in flesh. In all treatments, hue value of flesh and peel of ripe fruits were close to 90° which fall in the yellow quadrant. There is a significant difference in other parameters like fruit length, width and volume in 2 gl⁻¹ KCl-treated fruits when compared to control I indicating a significant increase.
Conclusion

Spraying of KCl at a concentration of 2 gl⁻¹ on mango fruits resulted in relatively high control of stem-end rot disease incidence and severity of ‘TomE JC’. Further, the level of residues of K was similar to untreated controls which may not cause harmful effects on the consumer. KCl treatment at 2 gl⁻¹ has a significant increase in yellowness of peel and flesh, fruit length, width and fruit volume. Thus, KCl can be used as an alternative, safe method in controlling stem-end rot in TomEJC mango.

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