Hydroponic Production of Purslane as a Sodium-removing Vegetable in NaCl-rich Nutrient Solution

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Abstract. To evaluate the potential of producing purslane (Portulaca oleracea L.) as a sodium (Na)–removing vegetable hydroponically at moderate NaCl salinity, two cultivars (Green and Golden) were grown in solutions with added 0, 6, 8, and 10 mM NaCl (the actual Na+ concentrations ≈ 2, 8, 10, and 12 mM, respectively). At harvest, 26 days after transplanting, apparent growth and biomass accumulation were not negatively affected by 6 to 10 mM added NaCl compared with 0 mM added NaCl. However, with the increase of added NaCl concentration from 0 to 6 to 10 mM, the sodium removal showed a 1- to 3-fold increase up to 0.26 to 0.41 mmol/plant, and 225.7 to 300.2 mmol·L−1 dry weight (DW) or 0.90 to 1.32 mmol·L−1·H2O, respectively. ‘Green’ produced greater biomass and removed more sodium per plant than ‘Golden’. ‘Golden’ had more of a dwarfed and compact canopy than ‘Green’. Sodium removal rate (mmol/plant/day) was the highest during the first 7 days after transplanting, but the fresh weight increase rate (g/plant/day) increased gradually as growth progressed. Results suggest that it is possible to hydroponically produce purslane in nutrient solutions with 8 to 12 mM Na+. Despite the high sodium-removal capability, purslane cannot be used to reduce Na+ concentrations in NaCl-rich hydroponic solutions. The biomass yield and the sodium removal of individual plants were affected by different cultivars and time after transplanting.

One of the major problems in closed hydroponic systems is the accumulation of salt ions, especially Na+ and Cl−, in reused solutions (Savvas et al., 2005). For most plant species, Na+ appears to reach a toxic concentration before Cl− (Munns and Tester, 2008), resulting in Na+ toxicity from a combination of osmotic and ionic stresses (Kronzucker and Britto, 2011). To prevent high Na+ levels in the rhizosphere, growers may periodically permit salt leaching (Varlagas et al., 2010). For example, in The Netherlands, discharge from closed hydroponic systems is allowed if Na+ concentration reaches 6 mM for cucumber or 8 mM for tomato during production (van Os, 1998). Discharging used nutrient solution can pollute the environment and sometimes even results in soil salinization (Varlagas et al., 2010). One way to address this problem is to use these solutions for the cultivation of another economically valuable crop, which has a high salt tolerance and the ability to accumulate sodium (Grieve and Suarez, 1997).

Purslane (Portulaca oleracea L.) is not only a salt-tolerant plant (Kilic et al., 2008; Shannon and Grieve, 1999), but also a good salt-removing crop (Aksoy et al., 2003; Grieve and Suarez, 1997), capable of removing up to 65 kg·ha−1 of Na+ in one growing season (Kilic et al., 2008). It has been recommended as a potential intercrop to remove salt in orchards (Kilic et al., 2008, 2010) and an excellent candidate for rehabilitation in the drainage water reuse systems (Grieve and Suarez, 1997). However, these studies have been carried out under soil or sand culture conditions (Grieve and Suarez, 1997; Kilic et al., 2008, 2010), and the related information is not available under hydroponic conditions, because Na+ uptake in plants can be affected by culture medium (Liu, 2002).

In addition, purslane is a traditional food crop in some Mediterranean, Central American, and Asian countries (Cros et al., 2007; Grieve and Suarez, 1997) and has now become a potential key vegetable crop worldwide because of its high nutritive and antioxidative properties (Uddin et al., 2012b; Wenzel et al., 1990). Recently, hydroponic production of purslane, as the easiest and cheapest growing method, has been gaining more and more attention as a result of the shorter cultivation cycles, higher planting densities, and clean and easily packed products compared with soil-based culture (Cros et al., 2007; Kaskar et al., 2008). Furthermore, previous studies indicated that shoots had considerably higher Na+ content than roots (Aksoy et al., 2003; Tester and Davenport, 2007) and, also, a tight coupling has been observed between plant growth (especially shoot growth) and Na+ uptake (Lv et al., 2012). So it may be possible to accumulate Na+ in purslane shoots at reasonably high levels without adversely affecting biomass productivity or, in other words, to hydroponically produce purslane as a sodium-removing vegetable.

Previous related studies in purslane were carried out separately on salt removal (Grieve and Suarez, 1997; Kilic et al., 2008) and vegetable production (Cros et al., 2007; Lara et al., 2011), and the NaCl concentrations in the solution used for growing purslane were totally different in these two categories of studies. In the salt removal studies (Grieve and Suarez, 1997; Kilic et al., 2008), the concentrations of NaCl treatment were usually higher up to more than 50 mM, but in the vegetable production studies (Cros et al., 2007; Lara et al., 2011), NaCl was not added to solutions and its concentrations were not known (should be very low). Currently, there is a lack of information regarding the effects of moderate NaCl concentrations (≥6 to 10 mM) on biomass production and sodium removal in purslane.

An appropriate selection of cultivars is vital to evaluate the potential of hydroponic production of purslane as a sodium-removing vegetable. Commercially available purslane cultivars are either green- or golden-leaved genotypes (Seedaholic, 2013). In studies aimed at evaluating growth and yield responses of purslane, significant differences were observed in plant growth between green- and golden-leaved genotypes in hydroponic solutions without added NaCl (Lara et al., 2011; Palaniswamy et al., 2000). However, in salt removal studies on purslane, only one genotype was used to evaluate Na+ removal capacity under high NaCl concentrations (Grieve and Suarez, 1997; Kilic et al., 2008). Consequently, comparisons of sodium removal and biomass accumulation between green- and golden-leaved purslane genotypes are needed under moderate NaCl concentrations (≥6 to 10 mM).

Determining how growth stage affects purslane biomass accumulation and sodium removal is important to predict the ideal harvest time of hydroponically produced purslane when used as a sodium-removing vegetable crop. A long production time (delaying the harvest time) would result in high biomass (i.e., increased yield); however, most crops exhibit higher Na+ uptake efficiency during vegetative vs. reproductive growth (Subbarao et al., 2003). Uddin et al. (2012a) found that Na+ concentration in dried purslane leaves decreased with plant maturity when evaluated over 60 d after transplanting young plants from the field into potted soil. In contrast, Kilic et al. (2008) reported that Na+ in dry purslane shoots was highest at the last harvest, but it was evaluated only 12 to 38 d after germination in a sand culture system. Therefore, further clarification is needed to evaluate how growth stages affect sodium removal and biomass accumulation of purslane.
hydroponically grown purslane during a longer growth period.

The objectives of the present study were to assess 1) the potential of producing purslane hydroponically under moderate NaCl concentrations (≈6 to 12 mM); 2) whether purslane can be used to remove sodium from hydropicnic solution with moderate NaCl salinity; and 3) how NaCl concentrations, cultivar type, and growth stage affect purslane growth and sodium removal from a hydropicnic system.

Materials and Methods

Plant materials and growing conditions.
The research was conducted in a greenhouse in the Edmund C. Bovey building at the University of Guelph, Guelph, Ontario, Canada (lat. 43°33’ N, long. 80°15’ W). Seeds of two purslane cultivars (Golden and Green; William Dam Seeds Co., Dundas, Ontario, Canada) were sown in rockwool cubes (1.5-inch Starter Plugs, Grodan Inc., Ontario, Canada) on 4 Jan. 2013. After germination, the seedlings were thinned to one plant per rockwool cube. Four plants (each in one rockwool cube), with two pairs of true leaves per plant, were transplanted, 26 d after sowing onto a Styrofoam disk floating on nutrient solution in a plastic pot (14-cm top and 12-cm bottom diameters × 15 cm high). Each pot contained ≈1.5 L of nutrient solution with macronutrients (mmol·L⁻¹) 19.9 nitrogen, 1.2 phosphorus, 10.8 potassium, 2.4 calcium, 0.1 magnesium, and 3.2 sulfur and micronutrients (μmol·L⁻¹) 8.3 copper, 19.1 boron, 9.6 manganese, 8.2 zinc, 1.7 molybdenum, 18.8 iron, and 0.4 EDTA. NaCl was added to the nutrient solutions to achieve four concentration treatments: 0, 6, 8, and 10 mM added NaCl. The initial Na⁺ concentration, electrical conductivity (EC), and pH values of the nutrient solutions are presented in Table 1. The nutrient solutions were changed every 7 d (except the last time, which was 5 d as a result of harvest) to reduce nutrient and NaCl concentration variability during the experiment. Nutrient solution pH ranged between 5.5 and 7.0 as measured at the end of each week before the solutions were renewed. Pots were arranged in a randomized block design with 10 blocks and eight treatments within each block (i.e., four NaCl concentrations for each of two purslane cultivars per block).

The greenhouse conditions were set at 18-h light/6-h dark by supplementing natural sunlight with high pressure sodium lamps to achieve a photosynthetic photon flux at canopy level averaging no less than 396.7 ± 33.5 μmol·m⁻²·s⁻¹, and 20 to 25 °C light/18/9 °C dark with a relative humidity between 60% to 80%.

Apparent growth measurements. One plant from each pot was randomly chosen from each block for growth measurements. Each week the following attributes were measured: main stem length, true leaf number on the main stem, side branch number on the main stem, and canopy width. Final harvest was carried out when the terminal flower bud cluster appeared on the main stem of both cultivars of purslane (26 d after transplanting).

Biomass accumulation estimation. Plant fresh weight (FW) increase during each week was determined by weighing the solutions and plants at the beginning and the end of each week, because the rockwool and Styrofoam weights were constant. Then, FW increase rate was calculated as follows:

\[ \text{FW increase rate (g/plant/d)} = \frac{(W_{ft} - W_{ips}) - (W_{it} - W_{ips})}{4 \times t} \]  

where \( W_{it} \) and \( W_{ft} \) are the final and initial total weight (g) of each pot together with solution, plants, rockwool cubes, and Styrofoam during each week, respectively. \( W_{ips} \) and \( W_{ips} \) are the final and initial weight (g) of each pot and solution during each week, respectively. \( t \) is the treatment time (d) of each week, and 4 is the number of seedlings in each pot.

At the time of harvest, each plant was cut at the base above the rockwool cube, and the remaining plant tissue was separated from the rockwool cube, washed with tap water, and then rinsed with deionized water. FW of plant tissue from above and within the rockwool cube were measured respectively. Plant tissue DWs were determined by drying in an oven at 65 °C until a constant weight was achieved.

 Marketable yield and water use efficiency were calculated as follows:

\[ \text{Marketable yield (kg·m}^{-2}\text{)} = \frac{FW_s}{1000 \times 0.015} \]  

where \( FW_s \) is the fresh weight (g) of harvested plant tissue above the rockwool cube from each pot and 0.015 is the top area (m²) of each pot.

Water use efficiency (g FW/L H₂O)

\[ \text{Water use efficiency} = \frac{\sum_{i=1}^{4} \Delta FW_i}{\sum_{i=1}^{4} \Delta V_i} \]

where \( \Delta FW_i \) is the increased fresh weight (g) of plants in each pot during the \( i \)th week (\( i = 1, 2, 3, 4 \)).

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Table 1. Initial Na⁺ concentrations, electrical conductivity (EC), and pH values of nutrient solutions used for hydroponic purslane production.

<table>
<thead>
<tr>
<th>Added NaCl concn (mM)</th>
<th>Measured Na⁺ concn (mM)</th>
<th>EC (dS·m⁻¹)</th>
<th>pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.8 ± 0.1</td>
<td>2.95 ± 0.10</td>
<td>6.9 ± 0.09</td>
</tr>
<tr>
<td>6</td>
<td>7.9 ± 0.1</td>
<td>3.59 ± 0.09</td>
<td>6.9 ± 0.08</td>
</tr>
<tr>
<td>8</td>
<td>9.9 ± 0.2</td>
<td>3.84 ± 0.09</td>
<td>6.9 ± 0.08</td>
</tr>
<tr>
<td>10</td>
<td>11.9 ± 0.3</td>
<td>4.04 ± 0.09</td>
<td>6.9 ± 0.08</td>
</tr>
</tbody>
</table>

aData are means ± se (n = 4).

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Fig. 1. Main stem length (A), number of true leaves (B), and number of side branches (C) on the main stem and canopy width (D) of ‘Green’ and ‘Golden’ purslane grown in hydropicnic solutions with different added NaCl concentrations at different growth stages. Data are means ± se (n = 10). Where bars are not visible, se does not exceed the size of the symbol. At each time point after transplanting, ns, *, **, or *** next to the data indicates that the effects of different cultivars (upper symbol) or NaCl concentrations (lower symbol) are not significant or significant at \( P \leq 0.05, 0.01, \) or 0.001, respectively.
Table 2. Biomass accumulation, marketable yield, and water use efficiency in ‘Green’ and ‘Golden’ purslane grown in hydroponic solutions with different added NaCl concentrations.

<table>
<thead>
<tr>
<th>Added NaCl conc (mM)</th>
<th>Fresh wt (g/plant)</th>
<th>Dry wt (g/plant)</th>
<th>Marketable yield (kg m⁻²)</th>
<th>Water use efficiency (g FW/L H₂O)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Green</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>22.1 ± 0.7a</td>
<td>1.30 ± 0.05 a</td>
<td>5.43 ± 0.18 a</td>
<td>80.2 ± 1.6 b</td>
</tr>
<tr>
<td>6</td>
<td>22.8 ± 0.6 a</td>
<td>1.33 ± 0.03 a</td>
<td>5.58 ± 0.14 a</td>
<td>85.6 ± 1.4 a</td>
</tr>
<tr>
<td>8</td>
<td>23.3 ± 0.9 a</td>
<td>1.33 ± 0.06 a</td>
<td>5.75 ± 0.22 a</td>
<td>86.8 ± 1.4 a</td>
</tr>
<tr>
<td>10</td>
<td>23.1 ± 0.9 a</td>
<td>1.39 ± 0.04 a</td>
<td>5.73 ± 0.21 a</td>
<td>84.9 ± 1.8 a</td>
</tr>
<tr>
<td><strong>Golden</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>21.2 ± 1.0 ab</td>
<td>1.13 ± 0.06 b</td>
<td>5.21 ± 0.26 b</td>
<td>79.6 ± 2.1 b</td>
</tr>
<tr>
<td>6</td>
<td>20.1 ± 1.1 b</td>
<td>1.06 ± 0.06 b</td>
<td>4.90 ± 0.26 b</td>
<td>83.2 ± 2.1 ab</td>
</tr>
<tr>
<td>8</td>
<td>20.7 ± 0.5 b</td>
<td>1.12 ± 0.03 b</td>
<td>5.08 ± 0.13 b</td>
<td>83.0 ± 1.6 ab</td>
</tr>
<tr>
<td>10</td>
<td>23.1 ± 1.1 a</td>
<td>1.32 ± 0.07 a</td>
<td>5.71 ± 0.27 a</td>
<td>84.1 ± 1.7 a</td>
</tr>
<tr>
<td><strong>ANOVA results</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G*</td>
<td>**</td>
<td>***</td>
<td>**</td>
<td>NS</td>
</tr>
<tr>
<td>C*</td>
<td>NS</td>
<td>**</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>G × C*</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
</tbody>
</table>

*Data are means ± se (n = 10).

2, 3, and 4), which can be estimated from Eq. (1), and ΔV is the volume (L) of consumed water in each pot during the ith week (i = 1, 2, 3, and 4). The initial and final volume of nutrient solution was measured weekly per pot and the difference in solution volume was assumed to be the weekly volume of water consumed by the four plants in each pot.

Sodium removal evaluation. A subsample of the fresh and week-old nutrient solution was analyzed for Na⁺ concentration using a portable pH/ISE meter and sodium electrode (Thermo Orion Model 290A, Waltham, MA). Na⁺ removal rate of the plants in each pot per week was calculated as follows:

\[
Na^+ \text{ removal rate (mmol/plant/d)} = \frac{V_i \times C_i - V_f \times C_f}{4 \times t}
\]

where \(V_i\) and \(V_f\) are the initial and final volumes (L) of solution in each pot during each week, respectively; and \(C_i\) and \(C_f\) are the initial and final Na⁺ concentrations (mM) in the solution in each pot during each week, respectively. \(t\) is the treatment time (d) of each week, and 4 is the number of seedlings in each pot. The Na⁺ uptake by rockwool and Styrofoam was subtracted by setting the blank controls (i.e., transplanting only rockwool and Styrofoam without plants into nutrient solution with different NaCl concentrations).

At harvest, total Na⁺ removal amount of each plant was calculated as follows:

\[
Na^+ \text{ removal amount (mmol/plant)} = \sum_{i=1}^{4} \text{SSR}_i \times t_i
\]

where \(\text{SSR}_i\) is the Na⁺ removal rate (mmol/plant/d) during the ith week (i = 1, 2, 3, and 4), which can be determined from Eq. (4), and \(t_i\) is the treatment time (d) during the ith week (i = 1, 2, 3, and 4). Eq. (6) was used for the calculation of total water consumption (L/plant) of each plant.

\[
\text{Water consumption (L/plant)} = \frac{\sum_{i=1}^{4} \Delta V_i}{4}
\]

where \(\Delta V_i\) is the volume (L) of consumed water in each pot during the ith week (i = 1, 2, 3, and 4), and the denominator in the formula, 4, is the number of seedlings in each pot.

Na⁺ removal efficiency (mmol·kg⁻¹ DW) and Na⁺ uptake concentration (mmol·L⁻¹ H₂O) were calculated as follows:

\[
\text{Na}^+ \text{ removal efficiency (mmol·kg⁻¹ DW)} = \frac{\text{SRA}}{\text{DW}} \times 1000
\]

\[
\text{Na}^+ \text{ uptake concentration (mmol·L⁻¹ H₂O)} = \frac{\text{SRA}}{\text{WC}}
\]

In Eqs. (7) and (8), SRA is the total Na⁺ removal amount per plant (mmol/plant), which is determined from Eq. (5). In Eq. (7), DW is the dry weight per plant (g/plant). In Eq. (8), WC is the total water consumption of each plant (L/plant), which can be determined from Eq. (6).
**Results**

**Apparent growth.** Apparent growth was not significantly affected by NaCl concentrations at any given time point during the 26 d of the study, but significant differences were obtained in most of the growth attributes between the two purslane cultivars (Fig. 1). The green-leafed cultivar had a longer main stem and wider canopy coverage than the golden-leafed one, which appeared to be more dwarfed and compact (Figs. 1A and 1D). Also, ‘Golden’ exhibited a shorter vegetative growth period than ‘Green’, because the terminal flower bud cluster appeared on 76.9% and 100% of the tested plants in ‘Golden’, but on only 0.6% and 38.8% of those in ‘Green’ at 16 and 26 d after transplanting, respectively.

**Biomass accumulation.** The rate of purslane biomass accumulation accelerated gradually after transplanting. The FW increase rate showed more than 2-fold increase during the trial, from 0.51 to 1.22 g/plant/d in ‘Green’ and from 0.44 to 1.26 g/plant/d in ‘Golden’, respectively (Fig. 2). NaCl significantly affected the FW increase rate in ‘Golden’ but not in ‘Green’. ‘Golden’ treated by 10 mM NaCl had a higher FW increase rate compared with 0 to 8 mM NaCl treatments 14 to 26 d after transplanting (Fig. 2B). On average, ‘Green’ exhibited a higher FW increase rate than ‘Golden’ from 0 to 21 d after transplanting.

At the time of harvest, the FWs and DWS and marketable yield were not significantly reduced by 6 to 10 mM NaCl treatments compared with 0 mM added NaCl (Table 2). Furthermore, the 10 mM NaCl treatment led to a higher DW of individual plants in ‘Golden’ and greater water use efficiency in both cultivars than the controls. On average, individual ‘Green’ plants had significantly higher FWs and DWSs and marketable yields than ‘Golden’, but no significant difference in water use efficiency.

**Sodium removal.** There were significant differences in sodium removal rates among different growth stages of purslane; each plant removed 2- to 4-fold greater mmol of sodium per day during 0 to 7 d than during 7 to 26 d after transplanting (Fig. 3). However, the significant cultivar differences were mainly observed during the latter stages rather than the former ones.

At final harvest, the amount of sodium removed by purslane increased linearly with the increase of Na+ as well as added NaCl concentrations in the nutrient solutions for both ‘Green’ and ‘Golden’ (Fig. 4A). Compared with 0 mM NaCl, plants treated with 6 to 10 mM NaCl removed more than 2- or 3-fold sodium (up to 0.24 to 0.41 mmol/plant) but consumed similar amounts of water per plant (Fig. 4A–B). Similarly, there were significant linear relationships between Na+ removal efficiency (mmol·kg⁻¹ DW), Na+ uptake concentration (mmol·L⁻¹ H₂O) and Na+ as well as added NaCl concentrations (Fig. 4C–D). On average, ‘Green’ showed significantly higher sodium removal, greater water consumption, and better Na+ uptake concentration than ‘Golden’ (Figs. 4A–B and 4D). However, no significant cultivar difference was obtained for Na+ removal efficiency (Fig. 4C).

**Discussion**

**Impacts of moderate NaCl salinity on hydroponic production and sodium removal of purslane.** Compared with 0 mM NaCl, 6 to 10 mM NaCl treatments did not inhibit apparent growth or reduce biomass accumulation and final yield in purslane in our study. However, reductions in plant growth caused by NaCl-dominated salinity were reported in several previous studies on purslane (Franco et al., 2011; Yazici et al., 2007). The discrepancy may be the result of differences in solution salinity levels; the EC values were less than 4.5 dS·m⁻¹ in our study but higher than 6.5 dS·m⁻¹, even up to 20 dS·m⁻¹ in previous studies. Purslane has been rated as a salinity-tolerant species with a yield threshold of 6.3 dS·m⁻¹ (Shannon and Grieve, 1999). Teixeira and Carvalho (2008) also showed that salinity-induced growth inhibition and biomass reduction were only observed in plants exposed to solutions with EC levels greater than 6.8 dS·m⁻¹ (≥60 mM NaCl).

In the present study, 10 mM added NaCl treatment (≥12 mM Na+ actual concentration) promoted dry biomass accumulation in ‘Golden’, suggesting sodium could be a beneficial element for this purslane cultivar. Recently, it has also reported that sodium, at reasonable concentrations, can promote biomass yield of some species including asparagus, barley, broccoli, carrot, pea, cabbage, kale, mustard, radish, celery, sugar beet, red beet, and turnip (Subbarao et al., 2003). For hydroponic production of purslane, it is clear that an addition of 6 to 10 mM NaCl to a solution (with an actual Na+ concentration of 8–12 mM) is not harmful, but rather beneficial for biomass accumulation, although for hydroponic production of most fruit vegetables (e.g., cucumber and tomato), these concentrations of Na+ may be damaging (Van Os, 1998, 1999).
Purslane was reported to preferentially absorb Na⁺ over other cations (e.g., K⁺, Ca²⁺, and Mg²⁺) with a 5- to 6-fold increase in shoot Na⁺ concentration when root zone salinity level increased from 2.1 to 28.5 dS·m⁻¹ (corresponding Na⁺ concentrations ranged from 15 to 275 mM; Grieve and Suarez, 1997). In the present study, when the added NaCl concentration increased from 0 to 6 to 10 mM (actual Na⁺ concentration increased from ≈2 to 8 to 12 mM) at a smaller interval, sodium removal by the purslane plants showed more than 2- or 3-fold increase. However, in fruit vegetables such as sweet pepper, less than 2- and 3-fold increase of Na⁺ accumulation (based on DW) was obtained in its leaves and fruits, respectively, when NaCl concentration in nutrient solution increased from 0 to 15 mM (actual Na⁺ concentration increased from ≈2 to 60 to 100 mM) (Carmassi et al., 2005). Obviously, purslane could remove far more Na⁺ from the nutrient solution with the same Na⁺ concentration, if consuming the same amount of water, as tomato. One fact to be kept in mind is that a large amount of sodium was removed from the solutions, which could reduce the total amount of sodium discharged to the environment; however, the solution sodium concentrations were not reduced by purslane in our study (data not shown).

Comparison of two purslane cultivars for hydroponic production and sodium removal. The ‘Green’ and ‘Golden’ did not exhibit any visual stress symptoms induced by 6- to 10-mM NaCl treatments indicating that both cultivars can tolerate these salinity levels. However, ‘Green’ exhibited more vigorous vegetative growth than ‘Golden’. ‘Green’ had a longer main stem and larger canopy. Palaniswamy et al. (2000) also found similar cultivar differences in plant growth with the green-leaved purslane being taller and having a larger leaf area per plant than the golden-leaved one grown in solution without added NaCl. Our study also showed that ‘Green’ had a longer vegetative growth period (as a result of a later appearance of flower buds) than ‘Golden’, indicating that ‘Green’ could be harvested as a leaf vegetable for a longer time.

Compared with ‘Golden’, ‘Green’ had a higher FW and DW per plant as well as a higher final marketable yield when grown at the same density. In non-NaCl-treated hydroponic solution, Palaniswamy et al. (2000) also reported that a green-leaved purslane showed higher FW and DW than a golden-leaved one. On the contrary, Lara et al. (2011) reported that a golden-leaved purslane cultivar had higher FW and DW than a green-leaved Spanish accession, C-215. Nevertheless, a common point in the two studies is that the purslane genotypes with higher biomass accumulation had a larger leaf area or a longer main stem regardless of leaf color. In another previous study, it was reported that fresh yield showed a positive linear relationship with main stem length, which was considered as a selection criteria for predicting purslane yield potential (Elmi et al., 1997). Thus, it seemed that the higher biomass accumulation and final yield of ‘Green’, in the present study, might not be associated with its leaf color, but with its longer main stem.

Significant cultivar differences also occurred in Na⁺ removal ability; ‘Green’ removed more Na⁺ per plant and showed a higher Na⁺ uptake concentration than ‘Golden’. There are several possible explanations for the differences: 1) it might be associated with a greater biomass accumulation in ‘Green’. A recent study on Salicornia europaea also indicated a tight coupling of biomass accumulation with sodium uptake (Lv et al., 2012); 2) it could be that ‘Green’ had a longer vegetative growth period than ‘Golden’, because plants translocate more Na⁺ to vegetative structures than to reproductive structures (Subbarao et al., 2003). ‘Green’ and ‘Golden’ had a similar sodium removal efficiency based on plant DW (mmol·kg⁻¹ DW). It may suggest that the two cultivars have similar NaCl salinity tolerance, because plant salt tolerance can be positively correlated with Na⁺ accumulation in plant tissues (Glenn et al., 1999).

Overall, it appeared that ‘Green’ had greater potential for hydroponic production as a vegetable than ‘Golden’ based on individual plant performance; based on a group of plants, ‘Golden’ may achieve higher biomass yield per unit area in the greenhouse through a higher planting density than ‘Green’. ‘Golden’ had more of a dwarfed and compact canopy than ‘Green’.

Effects of growth stage on biomass production and sodium removal of purslane. Our results suggest that purslane has a higher sodium removal rate in each plant during the first 7 d of growth than at the later growth stages, which is consistent with results reported by Uddin et al. (2012a), who reported a decrease in dry leaf concentration of Na⁺ as purslane plants matured. These facts indicate that purslane plants have the ability to control Na⁺ absorption and it could increase with time. A similar phenomenon was observed in wheat (Rivelli et al., 2002) and barley (Jeschke and Wolf, 1985; Rawson et al., 1988). The physiological basis for these responses is not known, although it is possible to be explained as follows. When a plant is exposed to a saline solution for the first time, a large amount of absorption of Na⁺ could provide a rapid means of turgor adjustment for plants to match the abrupt decrease in osmotic potential of the plant.
medium (Bernstein, 1975; Essah et al., 2003; Raven, 1985). Afterward, to avoid the toxicity of excessive Na+ accumulation, a plant could initiate many other osmotic adjustment ways such as synthesis of proline (Yazici et al., 2007), which would reduce Na+ uptake.

Unlike the pattern of Na+ absorption, the accumulation rate of fresh biomass in each plant increased gradually after transplanting, indicating that the yield efficiency of individual purslane plants could increase gradually with the delay of harvest time. It appeared that the period of maximum biomass accumulation rate did not match that of maximum sodium removal rate in individual plants. This contradiction implied that an earlier harvest (for example, 7 d after transplanting) of purslane plants would reduce the biomass accumulation to a large degree despite having the highest sodium removal rate in individual plants during this period and despite having the advantage of shortening crop production time.

It may be possible to compromise the contradiction derived from an earlier harvest at the level of group plants vs. individual plants by increasing the planting density, because the earlier the harvest time, the smaller the canopy width of individual plants (Fig. 1). Another possibility is to introduce larger purslane plants into NaCl-rich solutions only 1 week before harvest, which may maximize efficiency of both biomass accumulation and sodium removal. However, these speculations need further research for support.

In conclusion, it is possible to hydroponically produce purslane in nutrient solutions with 8 to 12 mM Na+. Although purslane has a higher sodium-removal capability than common fruit vegetables, it cannot be used to reduce Na+ concentration in NaCl-rich hydroponic solutions. The biomass yield and the sodium removal of individual purslane plants can be affected by different cultivars and time after transplanting.

**Literature Cited**


