Evaluating Growth and Yield Parameters of Five Quinoa (Chenopodium quinoa W.) Genotypes Under Different Salt Stress Conditions

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Abstract

Soil salinization is a global problem which restricts the choice of crop for cultivation. Management and reclamation of salinity using costly techniques may not be affordable by subsistence farmers. Therefore, it is important to look for new alternate crops like “quinoa” which are more salt-tolerant. As crops vary in their tolerance to salinity, they need to be evaluated for different salinity conditions. This study was conducted to evaluate five quinoa (Chenopodium quinoa W.) genotypes (ICBA-Q1, ICBA-Q2, ICBA-Q3, ICBA-Q4 and ICBA-Q5) for salinity tolerance under four artificially induced salinity (5, 10, 15, 20 dS m⁻¹) levels. The pot trials were conducted in a greenhouse, using 6 kg of Fluvisol soil in each pot. For comparison, trials were also conducted under field conditions. The parameters studied were rate of seed germination, plant height, fresh and dry biomass, chlorophyll content and grain yield. As expected, salinity had generally an inhibitory effect on all parameters. Out of the five quinoa varieties (ICBA-Q1 to ICBA-Q5), ICBA-Q3 and ICBA-Q4 proved to be more salt-tolerant. Therefore these two genotypes are recommended to farmers for large-scale adaptation.

Keywords: chlorophyll, germination, grain yield, quinoa, soil salinity, Ethiopia

1. Introduction

Increasing soil salinization has raised serious concerns of food security for the growing population of the world, which is expected to reach to 9 billion by 2050 (FAO, 2015). FAO estimated that over 1,000 million ha (mha) are globally affected by salinity and sodicity problems. Out of this, about 400 mha are saline, 450 mha are sodic and the remaining saline-sodic in nature (FAO, 2015). Currently, about 20% (62 mha) of the global irrigated land (over 300 mha) is affected with salinity whereas an additional 2,000 ha are added to this menace annually (Qadir et al., 2014). The growing existence of saline soils is reducing natural biodiversity as well as farm and livestock productivity. It is also threatening the sustainability of irrigated agriculture, which produces more than 80% of the total grains. Therefore there is a strong need to control spread of soil salinization. The possible solutions include using physical practices such as improved soil-water-crop management or adopt biological practices such as introducing salt-tolerant species (Ashenafi & Bob, 2016).

Ethiopia stands first in Africa in the extent of salt-affected soils with an estimated 11 mha of land exposed to salinity (Ashenafi & Bob, 2016; Frew et al., 2015). This relates to 13% of the total irrigated area of the country (Birhane, 2017). The saline soils are mainly located in the Rift Valley, Wabi Shebelle River Basin, the Denakil Plains and other lowlands of the country, where about 10% of the population lives (Sileshi, 2016). The problems of soil salinity are expected to increase in future due to the establishment of large-scale irrigation schemes without the provision of adequate drainage facilities (Birhane, 2017). The salinity problems have grave implications for the future food security and economic development of the country. With an annual population growth rate of 3%, securing food and improving livelihood of the rising population will be the biggest challenge. Even today, food shortages are widespread and since 1970 country is in the grip of consistent famines. It is reported that among children aged up to 5 years, around 25% are underweight and 40% are stunted due to malnutrition (UNCEF, 2016).

In the Rift valley and other irrigated areas of Ethiopia, salinity development is mainly caused by the presence of soluble salts in the irrigation water, hot and dry weather conditions and excessive use of poor quality
groundwater for irrigation. Development of large irrigation schemes in middle and lower Awash Valley without effective drainage systems along with poor water management practices have resulted in the gradual rise of saline groundwater. Due to high evapotranspiration, water evaporates from the soil surface leaving the salt behind causing secondary salinization in these areas (Frew et al., 2015). The farmers in Ethiopia are mainly using flood/basin methods to irrigate their poorly levelled fields. There is a tendency to over-irrigate because farmers usually do not have enough knowledge of actual crop water requirements. Therefore, if the current irrigation practices will continue, salinity problems will further exacerbate. Therefore there is an urgent need to take necessary measures to control spread of soil salinization.

In order to meet the future food security challenges, reclamation of existing saline soils and prevention of other areas from salinity development is of paramount importance. The low to medium salinity areas can be reclaimed through effective leaching and installation of appropriate drainage systems. However, these strategies are costly, time consuming and difficult to implement by farmers due to lack of financial resources and technical know-how (Qadir & Oster, 2004). The highly saline soils can be reclaimed by using chemical amendments and/or adoption of biosaline approach. The biosaline approach entails introduction of salt-tolerant food and forage crops, which can withstand higher salinity levels. These integrated food and forage systems can help in increasing flexibility of smallholder farmers to feed their families and livestock. However, this approach requires selection of diverse food and forage species which have the capacity to resist salts present in the soil (Qureshi, 2017).

One such crop is Quinoa (Chenopodium quinoa Willd), which has emerged as an ideal crop for drought prone and salinized agricultural areas due to its high nutritious value and wide adaptability and ability to grow in harsh climatic conditions characterized by high temperatures and poor soil and water quality (Ruiz et al., 2014; Bazile et al., 2016; Chukar-Allah et al., 2016). Quinoa is grown in Andes region for centuries, however, its production and consumption outside the Andes is relatively new (Jacobsen, 2017). Currently, quinoa is grown in more than 90 countries. Today, 80 percent of the quinoa production comes from Bolivia and Peru whereas the remaining 20 percent is produced in all other countries (Bazile et al., 2016). Despite this rapid exposition, quinoa farming is still in “experimental phase” in many countries. The yield differences are huge ranging from 0.6 to 3.9 t ha \(^{-1}\) depending on soil, water and climatic conditions (Scalin & Lewis, 2017). This clearly indicates the need for further research to develop varieties that can produce consistent yields under different agro-climatic conditions.

Quinoa has successfully been grown from non-saline to highly saline soils (15-20 dS m \(^{-1}\)) (Wilson et al., 2002; Bosque Sanchez, 2003; Adolf et al., 2012). At these salt levels other plant types either fail to grow or grow only poorly (Munns & Tester, 2008; Shabala et al., 2013)). Besides being gluten free, quinoa grain is rich in proteins and essential amino acids such as lysine, threonine, methionine, and much needed unsaturated fatty acids (i.e., linoleic, oleic and linolenic), of minerals (Ca, Fe, Cu, Zn) and vitamins (A, B2, C and E) (Vego-Gálvez et al., 2010). Quinoa is a good source of calcium and is suitable for lactose-intolerant consumers and those allergic to gluten (Repo-Carrasco et al., 2003; Vega-Gálvez et al., 2010). The leaves of the plant are frequently eaten as a leafy green vegetable just like as spinach. It can also be used as a highly nutritious feed for animals.

Quinoa is new in African countries, currently passing through pilot testing and field trial stages. Quinoa could withstand temperatures from -8 °C to 38 °C, at sea level or 4,000 meters above, which makes it viable for farmers and the crop still in “experimental phase” in many countries. The yield differences are huge ranging from 0.6 to 3.9 t ha \(^{-1}\) depending on soil, water and climatic conditions (Scalin & Lewis, 2017). This clearly indicates the need for further research to develop varieties that can produce consistent yields under different agro-climatic conditions.

2. Materials and Methods

2.1 Study Area

The experiments were conducted at the Werer Agricultural Research Center (WARC), Amibara, Ethiopia, which is located at 278 km to the east of Addis Ababa (9°12′18″ N latitude and 40°15′21″ E longitude). The study area is relatively flat with slope gradients of 1-2% (Figure 1). The mean annual rainfall is 570 mm with a minimum and maximum temperatures of 19 °C and 34 °C, respectively (Figure 2). Higher soil evaporation due to extreme temperatures causes the creation of saline soils and nutrient disparity in the soils causing poor plant growth. The Vertisols soil type of the area varies from silty clay to clay whereas the texture of the Fluvisols soils sandy loam to silty loam (Heluf, 1985; Wondimagegne & Mnalku, 2012).
2.2 Trials Under Controlled Conditions

For pot trials under controlled conditions, treatments include two factors; food and forage genotypes and salt stress levels. The trials were conducted under 4 salt stress conditions to evaluate its suitability for different regions. Four salt stress treatments were prepared by mixing 7.28 14.57, 21.43 and 29.14 g of NaCl into 6.0 kg of soil packed per pot to produce salinity levels of 5, 10, 15 and 20 dS m⁻¹. Five quinoa genotypes (ICBA-Q1, ICBA-Q2, ICBA-Q3, ICBA-Q4, ICBA-Q5) were evaluated to test their performance under different soil salinity conditions. The treatments were organized in a completely randomized design with three replications.

Ten seeds of each genotype were sown in each pot. The seeds were surface sterilized using 70% ethanol (exposure for 10 seconds) followed by immersion for 10 minutes in sodium hypochlorite solution (NaClO; 5% active chloride). The treated seeds were washed thoroughly with distilled water and were placed on moist filter paper in petri dishes. Uniformity of seed size and quality was ensured before germination test. Irrigations were done with canal water (EC = 0.3 dS m⁻¹).

2.3 Trials Under Field Conditions

Field trials were conducted on the saline-sodic soil with an average ECe value of 19.50 dS m⁻¹ and an exchangeable sodium percentage (ESP) value of 20. A plot size of 3 m × 4 m was used for field experiments. Soil samples were collected from 0-30 cm depth and analyzed for different soil parameters. The soil bulk density was determined according to the method described by Black (1965). The exchangeable bases (Ca, Mg, Na, and K) were determined from the extraction of neutral normal ammonium acetate extraction, Ca and Mg from EDTA titration method, while K and Na using flame photometer. The cation exchange capacity (CEC) of soil was
determined by the percolation tube procedure (Van Reewijk, 1992). The ESP was computed as the percentage of the exchangeable Na to the CEC of the soil (Table 1).

Table 1. Soil physio-chemical properties of the field trial site

<table>
<thead>
<tr>
<th>Quinoa genotypes</th>
<th>Soil depth (cm)</th>
<th>pH</th>
<th>ECe (dS m⁻¹)</th>
<th>Exchangeable bases (cmol(+), kg⁻¹)</th>
<th>CEC (cmol(+), kg⁻¹)</th>
<th>ESP (%)</th>
<th>BD (g cm⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICBA-Q1</td>
<td>0-30</td>
<td>7.9</td>
<td>19.32</td>
<td>Ca+Mg: 36.94, Na: 8.00, K: 1.04</td>
<td>41.16</td>
<td>19.44</td>
<td>1.38</td>
</tr>
<tr>
<td>ICBA-Q2</td>
<td>0-30</td>
<td>7.8</td>
<td>20.34</td>
<td>Ca+Mg: 43.02, Na: 7.68, K: 0.96</td>
<td>44.21</td>
<td>17.37</td>
<td>1.36</td>
</tr>
<tr>
<td>ICBA-Q3</td>
<td>0-30</td>
<td>8.0</td>
<td>20.01</td>
<td>Ca+Mg: 38.83, Na: 9.67, K: 1.01</td>
<td>39.83</td>
<td>21.28</td>
<td>1.37</td>
</tr>
<tr>
<td>ICBA-Q4</td>
<td>0-30</td>
<td>8.1</td>
<td>18.76</td>
<td>Ca+Mg: 37.91, Na: 9.01, K: 0.98</td>
<td>41.78</td>
<td>21.57</td>
<td>1.39</td>
</tr>
<tr>
<td>ICBA-Q5</td>
<td>0-30</td>
<td>7.9</td>
<td>18.98</td>
<td>Ca+Mg: 38.02, Na: 8.98, K: 0.87</td>
<td>43.57</td>
<td>20.61</td>
<td>1.36</td>
</tr>
</tbody>
</table>

Since soils of the study area are good in nutrients therefore no fertilizer was used for experiments. Irrigations were given according to crop evapotranspiration (ETc), which was calculated by multiplying reference evapotranspiration (ETo) with the crop coefficient (Kc). ETo was calculated using modified Penman-Monteith equation whereas the Kc values were taken from FAO-56 (Allen et al., 2006). In addition to total irrigation requirements according to ETc, an additional 10% of the total irrigation amount was applied for leaching of salts to manage root zone salinity.

2.4 Observations and Measurements

Mean germination time (MGT), germination percentage (GP), biomass and grain yield and shoot and root dry matter and other related data was measured. Seeds with full radicle were considered as germinated. Germination count was done on 5th, 10th and 15th day after plantation. GP was calculated according to Ashraf and Foolad (2005) whereas MGT was determined using equation of Ellis and Roberts (1981). Chlorophyll content (SPAD units) of leaves was measured using Minolta Soil-Plant-Analysis Development (SPAD) meter. Plant height was measured with a standard ruler (i.e., stem length from soil level to the top of the flower head).

\[ GP = \frac{\text{Total germinated seeds}}{\text{Total number of seeds}} \]  
\[ \text{MGT} = \frac{\sum Dn}{\sum n} \]

Where, \( n \) = Number of germinated seeds on day \( D \), and \( D = \text{Number of days from the start of germination}. \)

2.5 Statistical Analysis

Field and pot experiments were conducted for two years (2017-18) and the data was subjected to analysis of variance (ANOVA) technique (A. Gomez & H. Gomez, 1984) for factorial CRD using SAS 9.3 software (SAS Institute, Cary, NC). The significance of differences between the mean values at \( p < 0.05 \) was determined using Least Significance Difference (LCD) test. The comparison between all data obtained was made by using Duncan’s Multiple Range Test (DMRT).

3. Results

3.1 Trials Under Controlled Conditions

3.1.1 Germination Percentage (GP), Mean Germination Time (MGT) and Germination Index (GI)

For all genotypes, increasing salinity affected seed germination. The GP was highest in ICBA-Q3, ICBA-Q4 and ICBA-Q5 in control and gradually decreased with the increasing salinity. The lowest GP was found in ICBA-Q1 and ICBA-Q2. The MGT also increased with growing salinity levels. The highest MGT was recorded in ICBA-Q1 at 20 dS m⁻¹, followed by ICBA-Q2 whereas the lowest was found in ICBA-Q3 in control. MGT for other three genotypes were comparable (Table 2). The GI followed the trends of GP for all genotypes. The maximum GI was observed in ICBA-Q5 followed by ICBA-Q3 and ICBA-Q4 at control. Lower GI values were observed at the highest salt concentration levels for all genotypes.
Table 2. Effects of salinity on GP, MGT and GI of five quinoa genotypes

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Genotypes</th>
<th>NaCl salt level (dS m⁻¹)</th>
<th>LSD (p ≤ 0.05)</th>
<th>CV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>5</td>
<td>10</td>
</tr>
<tr>
<td>Germination Percentage (%)</td>
<td>ICBA-Q1</td>
<td>36.67</td>
<td>23.33</td>
<td>10.00</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q2</td>
<td>16.67</td>
<td>16.67</td>
<td>13.33</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q3</td>
<td>83.33</td>
<td>80.00</td>
<td>63.33</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q4</td>
<td>83.33</td>
<td>83.33</td>
<td>56.67</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q5</td>
<td>83.33</td>
<td>86.67</td>
<td>63.33</td>
</tr>
<tr>
<td>Mean Germination Time (days)</td>
<td>ICBA-Q1</td>
<td>2.67</td>
<td>3.27</td>
<td>5.88</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q2</td>
<td>3.11</td>
<td>3.33</td>
<td>5.66</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q3</td>
<td>2.61</td>
<td>3.83</td>
<td>5.27</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q4</td>
<td>3.00</td>
<td>4.33</td>
<td>5.61</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q5</td>
<td>3.16</td>
<td>4.33</td>
<td>5.67</td>
</tr>
<tr>
<td>Germination Index (GI)</td>
<td>ICBA-Q1</td>
<td>1.01</td>
<td>0.45</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q2</td>
<td>0.31</td>
<td>0.21</td>
<td>0.19</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q3</td>
<td>2.38</td>
<td>2.17</td>
<td>1.57</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q4</td>
<td>2.02</td>
<td>2.32</td>
<td>2.01</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q5</td>
<td>2.59</td>
<td>2.51</td>
<td>1.92</td>
</tr>
</tbody>
</table>

3.1.2 Plant Height

For all quinoa genotypes, a decreasing trend in plant height was observed with the increasing level of salinity. The highest plant height was observed for ICBA-Q3 (92.7 cm) and ICBA-Q4 (92.3 cm) at 0 dS m⁻¹. However, at higher salinity level (20 dS m⁻¹), plant heights of ICBA-Q3 and ICBA-Q4 were reduced to 54.6 cm and 51.0 cm, respectively. Increase in salinity from 0 to 20 dS m⁻¹ causes reduction in plant height of for ICBA-Q3 and ICBA-Q4 by 41% and 44%, respectively. Table 3 shows that plant height of all quinoa genotypes reduced significantly after 10 dS m⁻¹. These results agree with those of Jacobsen (2003) and Al-Dakheel et al. (2015) who found significant reduction in plant height with the increasing salinity levels in Phaseolus species and Lentils.

Table 3. Effects of salinity on plant height of five quinoa genotypes

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Genotypes</th>
<th>NaCl salt level (dS m⁻¹)</th>
<th>LSD (p ≤ 0.05)</th>
<th>CV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plant height (cm)</td>
<td>ICBA-Q1</td>
<td>68.67</td>
<td>67.00</td>
<td>67.67</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q2</td>
<td>68.33</td>
<td>67.67</td>
<td>67.33</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q3</td>
<td>92.67</td>
<td>84.00</td>
<td>74.67</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q4</td>
<td>92.33</td>
<td>85.00</td>
<td>69.47</td>
</tr>
<tr>
<td></td>
<td>ICBA-Q5</td>
<td>70.00</td>
<td>65.33</td>
<td>60.67</td>
</tr>
</tbody>
</table>

3.1.3 Dry Biomass Yield

Dry biomass yield was attained by oven-drying fresh biomass at 65 °C to constant weight. In all quinoa genotypes, dry biomass yield conceded due to increased salt stress (Figure 3). The highest dry biomass yield was obtained in ICBA-Q3 at 0-5 dS m⁻¹ whereas the lowest was obtained in ICBA-Q1 and ICBA-Q2. ICBA-Q4 performed better at elevated salinity levels (15-20 dS m⁻¹). The dry biomass yield decreased with the increasing soil salinity in the growth medium, although the response of all five genotypes to different salinity levels was heterogeneous. At 0 dS m⁻¹, dry biomass yield of ICBA-Q1, ICBA-Q2, ICBA-Q3, ICBA-Q4, and ICBA-Q5 was 15.5, 12.5, 29.6, 25.0 and 21.0 g/plant, respectively. However, the dry biomass yield at 20 dS m⁻¹ was noted as 5.1, 8.6, 17.1, 18.1, and 13.5 g/plant, registering a reduction of 67%, 30%, 42%, 28%, 36% for ICBA-Q1, ICBA-Q2, ICBA-Q3, ICBA-Q4, and ICBA-Q5, respectively. For all salinity levels, minimum reduction was noted in ICBA-Q3, which shows that this genotype is most resistant to increasing salinity levels.
Figure 3. Dry biomass yield of five quinoa genotypes as affected by different salinity levels

The experimental data was also used to develop production functions for 5 quinoa genotypes under different soil salinity levels and the results are presented in Figure 4. The highest reduction in dry biomass yield per unit increase in salinity (1 dS m\(^{-1}\)) was observed in ICBA-Q3 (0.56 g/plant) followed by ICBA-Q1 (0.48 g/plant) and ICBA-Q2 (0.46 g/plant). The lowest reduction in dry biomass per unit increase in salinity was found in ICBA-Q4 (0.38 g/plant) and ICBA-Q5 (0.22 g/plant). These two genotypes showed more stable dry biomass yields under all salinity levels. The dry biomass yield for ICBA-Q3 sharply declined after 5 dS m\(^{-1}\) salinity level whereas this was not the case for ICBA-Q4 and ICBA-Q5. ICBA-Q5 showed more consistent dry biomass yield at all salinity levels. This shows that under moderate salinity levels (0-5 dS m\(^{-1}\)), ICBA-Q3 can be a good choice because of high yielding potential. However, for higher salinity levels (10-20 dS m\(^{-1}\)), ICBA-Q4 and ICBA-Q5 are more suitable due to their higher salt tolerance capacity.
3.1.4 Grain Yield

The grain yield was negatively affected by increasing salinity levels however, the impact was more noticeable at higher salt concentrations (Figure 5). Since ICBA-Q1 and ICBA-Q2 were poor in germination, they also produced lower grain yield. The highest grain yield was obtained for ICBA-Q3 at all salinity levels followed by ICBA-Q4 and ICBA-Q5. The differences in grain yields under ICBA-Q3 and ICBA-Q4 were non-significant. Under control, the grain yield of ICBA-Q4 and ICBA-Q5 was 10% and 42.5% less than the grain yield of ICBA-Q3. However, the reductions in grain yields at the higher salinity levels were comparatively lower than the control i.e., grain yields of ICBA-Q4 and ICBA-Q5 were 4.8% and 38% less than ICBA-Q3. The grain yields of ICBA-Q3 and ICBA-Q4 were comparable at 10-15 dS m⁻¹, however, there was a significant reduction in grain yields at higher salinity level (20 dS m⁻¹).

Figure 6 shows that the maximum reduction in grain yield per unit increase of salinity was recorded for ICBA-Q3 genotype followed by ICBA-Q4 genotype. Although ICBA-Q1 and ICBA-Q2 genotypes produced lower yields, the reduction in grain yield per unit increase in salinity for these genotypes was lower, e.g., 0.28 g/plant for ICBA-Q1 and 0.21 g/plant for ICBA-Q2. The yield reduction per unit increase in salinity in ICBA-Q5 was 0.35 g/plant. Considering the overall dry biomass yield, grain yield and tolerance against higher salinity levels, ICBA-Q3 and ICBA-Q4 can be considered as the best genotype for all salinity levels under Ethiopian conditions.
3.1.5 Chlorophyll Content

Due to the distinct variation among all five quinoa genotypes, their responses to different salinity regimes for chlorophyll content were also different. Figure 7 illustrates that the chlorophyll content of all five genotypes tolerated salinity stress up to 10 dS m⁻¹ but decreased significantly at higher salinity levels. The highest chlorophyll content was recorded at salinity levels of 0-5 dS m⁻¹. The results indicate a decreasing trend of chlorophyll content with increasing salinity stress except in ICBA-Q2, ICBA-Q4, and ICBA-Q5 where chlorophyll content was higher at 5 dS m⁻¹ compared to control. At lower salinity levels (0-5 dS m⁻¹), the performance of ICBA-Q4 and ICBA-Q5 was superior than other three genotypes. Net decrease in the photosynthesis rates of quinoa by high salinity has also been reported (Eisa et al., 2017). They have also reported salt-induced growth reduction due to low photosynthetic supply because of impaired photosynthetic capacity.
For the validation of pot trials results, selected five quinoa genotypes were also tested under field conditions with soil salinity (ECe) values approaching 20 dS m\(^{-1}\) (Table 4). The tested quinoa genotypes were found to be significantly different in biophysical parameters such as grain yield, days of 50% physiological maturity, days of 50% emergency, number of panicles per plant, plant height, number of days to pasty grain and milky stage and dry biomass yield. The results indicate that ICBA-Q3 performed superior with regard to grain and dry biomass yield followed by ICBA-Q4. The performance of ICBA-Q1 and ICBA-Q2 was the poorest under field conditions whereas ICBA-Q4 and ICBA-Q5 showed medium results. The field evaluation shows that ICBA-Q3 is the most suitable genotype for hot, dry and highly saline areas both in terms of dry biomass and the grain yield. As evident from the pot trial analysis, other genotypes such as ICBA-Q4 and ICBA-Q5 can yield satisfactory results in low to medium saline areas. Therefore, quinoa genotype for a certain area should be selected after due consideration of climatic and soil salinity conditions.

### Table 4. Field evaluation of five salt-tolerant quinoa genotypes

<table>
<thead>
<tr>
<th>Quinoa genotypes</th>
<th>DE (days)</th>
<th>DM (days)</th>
<th>PH (cm)</th>
<th>PPP (days)</th>
<th>DMS (days)</th>
<th>DPG (days)</th>
<th>DBY (ton/ha)</th>
<th>GY (kg/ha)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICBA-Q1</td>
<td>12.33</td>
<td>93.00</td>
<td>138.60</td>
<td>8.00</td>
<td>80.00</td>
<td>90.67</td>
<td>1.239</td>
<td>464</td>
</tr>
<tr>
<td>ICBA-Q2</td>
<td>12.00</td>
<td>94.33</td>
<td>148.77</td>
<td>8.00</td>
<td>78.33</td>
<td>89.67</td>
<td>1.291</td>
<td>499</td>
</tr>
<tr>
<td>ICBA-Q3</td>
<td>9.00</td>
<td>86.33</td>
<td>144.00</td>
<td>11.67</td>
<td>72.33</td>
<td>84.00</td>
<td>7.211</td>
<td>2965</td>
</tr>
<tr>
<td>ICBA-Q4</td>
<td>8.67</td>
<td>89.00</td>
<td>152.13</td>
<td>10.00</td>
<td>75.67</td>
<td>88.00</td>
<td>5.885</td>
<td>1644</td>
</tr>
<tr>
<td>ICBA-Q5</td>
<td>8.67</td>
<td>86.00</td>
<td>156.13</td>
<td>9.00</td>
<td>71.33</td>
<td>78.66</td>
<td>4.023</td>
<td>1559</td>
</tr>
<tr>
<td>LSD (P &lt; 0.05)</td>
<td>1.49</td>
<td>3.94</td>
<td>NS</td>
<td>1.41</td>
<td>NS</td>
<td>6.68</td>
<td>0.572</td>
<td>152</td>
</tr>
</tbody>
</table>

**Note.** DE = Days to 50% Emergence; DM = Days to 50% Maturity; PH = Plant Height; PPP = Panicles per Plant; DMS = Days to Milky Stage; DPG = Days to Pasty Grain; DBY = Dry Biomass Yield; GY = Grain Yield.

### 4. Discussion

The rising global demand for nutritious and healthy food has challenged scientists to look for alternate crops especially for the marginal areas where agricultural production is inefficient due to unfavorable climatic conditions, low soil fertility and lack of good quality irrigation water. In many countries of the Middle East and Africa region, scientists are experimenting quinoa production because it is rich in nutrients, tolerant to salinity and uses much less water than other crops. Against this backdrop, this study was focused on assessing the feasibility of 5 different quinoa genotypes for dry and saline soil conditions of Ethiopia under controlled and field conditions. The seed germination was adversely affected by the rising salinity. The salinity impedes seed germination either without loss of viability at higher salinities and/or by inducing stress to seeds (Breusegen et al., 2006). Gómez-Pando et al. (2010) did a study on 15 most salt-tolerant Peruvian accessions of quinoa and found that some genotypes of quinoa showed decline in germination and plant height under high saline conditions, while others did not or even register an increase.
The results indicate reduction in dry biomass yield with increasing salinity, which might be due to lack of water availability and hydrolysis of reserved foods and their translocation to the growing shoots. Other factors responsible for lower dry biomass yield may include panicle length, chlorophyll concentrations, number of productive tillers, number of primary branches per panicle, and fertility percentage (Ali et al., 2004). Studies have also reported reduction in plant growth and dry-matter accumulation under saline conditions in several grain legumes including *P. vulgaris* that can be ascribed by decrease in cell elongation (Turan et al., 2007; Gómez-Pando et al., 2010).

Gómez-Pando et al. (2010) have also found a remarkable influence of quinoa genotypes on root dry mass per plant under saline conditions. This was probably due to the stunted growth of plants caused by high salt concentration in the nutrient medium. The higher salt stress causes reduction in the rate of leaf surface expansion, which results in considerable decrease in the dry weights of shoot, leaves, and roots (Kandil et al., 2012). This can be linked to the limited supply of metabolites to young growing tissues. Metabolic production usually occur within the leaves and can be affected significantly at high salt stress conditions either due to the low water uptake or toxic effect of NaCl concentration (Hassen et al., 2014).

The results of this study also show decrease in chlorophyll content with the increasing salinity levels in all tested quinoa genotypes. This can be attributed to the fact that at the higher salinity levels, plants increase generation of reactive oxygen species as by-product. This damages the cellular components and cause chlorophyll degradation and membrane lipid peroxidation, reducing membrane fluidity and selectivity (Verma & Mishra, 2005). Reduction in chlorophyll content due to metabolic limitations of photosynthesis in leaves at higher salinity levels (above 250 mM) has also been reported by Munns et al. (2006). In another study decreasing chlorophyll content in trees is associated with aggravated salt stress due to enzymatic chlorophyll degradation (Xu et al., 2011).

The reduction in the photosynthesis under saline conditions mainly occur due to a reduction in leaf area, chlorophyll content and stomatal conductance, and decrease in photosystem II efficiency. The reduction in chlorophyll levels in plants under saline conditions is considered as a typical symptom of oxidative stress and is usually linked with the lack of chlorophyll synthesis, together with the activation of its degradation by the enzyme chlorophyllase (Netondo et al., 2004). The decrease in chlorophyll results in reduced biomass with respect to increasing salinity stress.

The decreases in chlorophyll with increased salt stress were found in *Phaseolus vulgaris* L., and cereals (Santos, 2004). Cocozza et al. (2013) has also shown that quinoa plant is more resistant to water and salt stresses due to its effective stomatal responses that helps plant growth and protected crop yield (Sai Ram et al., 2002). Jaleel et al. (2008) has shown reductions of chlorophyll content in *Catharanthus roseus* whereas Nazarbeygi et al. (2011) has got similar results in Canova. Accumulation of salts causes an irreparable damage to the photosynthetic mechanism due to dehydration of cell membranes and closure of stomata which reduces their permeability to CO₂ and thus chlorophyll formation.

5. Conclusions

There are considerable differences on various plant growth parameters with the increasing salinity on five quinoa genotypes. Results clearly revealed that nearly all parameters measured decreased with increasing levels of salinity stress. Most limiting factor for decreased plant growth was found to be reduction in photosynthesis expressed in the production of chlorophyll. We suggest that in future, plant breeding should focus on developing new genotypes that can withstand salinity and have high antioxidant activity. This study has shown that the performance of ICBA-Q3 is superior followed by ICBA-Q4 and ICBA-Q5. However, further optimization of these genotypes is recommended to enhance their productivity under different agro-ecological conditions. This should include testing of these genotypes for water and heat stresses to assess their suitability for hot and dry climatic conditions prevailing in many regions of Ethiopia. In this study, fresh water is used for all experiments. However, in many areas, availability of fresh water is limited, and poor quality groundwater is used for irrigation. Therefore, it would be worthwhile to evaluate the performance of these quinoa genotypes using different quality irrigation waters.

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