Presoaking Treatment of Propolis Aqueous Extract Alleviates Salinity Stress in Spinach (Spinacia oleracea L.) Plants Grown under Calcareous Saline Soil Conditions

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Abstract. Two pot experiments were conducted during the two successive seasons of 2014 and 2015 to study the effect of propolis extract at the rates 0, 6000, 7000, 8000 and 9000 mg/L solution used as seed soaking to spinach seedlings on growth, yield and some chemical constituents of spinach plants (Spinacia oleracea L.) grown under calcareous saline soil conditions. The obtained results indicated that increasing the rates of propolis extract as seed soaking application increased the growth parameters of the treated plants. The best result was obtained by the middle rate (7000 mg/L) as seed soaking in both seasons of the study. The same trend was also observed regarding all studied chemical constituents, i.e. chlorophyll a, b and total carotenoids concentration, anthocyanine, total carbohydrates, total and reducing sugars, total free amino acid, free proline, crude protein, total indoles, total phenols, N, P and K in leaves. Moreover, soaking seeds in propolis extract before planting improved the metabolic activity of seeds through the increase in seed values in total and reducing sugars, total free amino acid, total indoles and total phenols as well as the lowest values of total carbohydrate. Thus, the coincident application of propolis extract at (7000 mg/L) as a seed soaking ingredient is recommended for improving growth, yield and chemical composition of spinach plants and for overcoming the adverse effect of saline conditions.

1. Introduction

Spinach could be a common vegetable within the people's everyday life mainly for its characteristic inexperienced color, organic process content like carotenes, vitamin C, and minerals like calcium and iron [1]. The intake of a specific quantity of vegetables will facilitate reducing the risk of many diseases like cancer and various disease [2]. Cultivated globally, spinach is additionally a very important raw-material within the food process industry [3]. A general reduction in growth and yield because of salinity is widely documented [4, 5] on Spinach plants that, growth and yield of spinach plants were minimized by increasing soil salinity. Many investigators have conducted studies for up salt tolerance of plants [5-7].

In recent years, there is a growing interest with natural bio-stimulating substances. Propolis (bee glue) is the generic name for the pitchy substance collected by honeybees (Apis mellifera L.) and found to be effective against a spread of microorganism, bacteria, viruses and fungi. There are varied literature knowledge of characteristically the chemical composition of propolis [8-13]. Propolis contains many necessary compounds has been detected to have an effect on the activity of many physiological processes in plants [8, 13]. Amino acids, sugars, bound vitamins (particularly, B-group, C and E), minerals, terpenes and sesquiterpenes are considered or thought-about to be among these necessary compounds. We are aware that terpenes and sesquiterpenes can be significant compounds important to plant growth processes. Terpenoids are thought-about to be the precursors of the many phytohormones (particularly, gibberellins), that are necessary for plants grown under numerous stresses.

The useful impact of propolis extract on growth, yield and chemical constituents of plants was accordingly on several species of plants [14-16, 7, 17].
Accordingly, the aim of this work was to check the impact of propolis extract as seed presoaking agent on growth, yield and chemical composition of mature spinach plants under saline conditions of saline calcareous soil and to clarify the role of propolis extract in minimizing the injurious impact of salinity on spinach plants.

2. Materials and Methods

The present investigation was conducted during the two successive seasons 2014 and 2015 in the Experimental Station, Faculty of Agriculture, Fayoum University, Egypt. The physical and chemical properties of the soil were tested by the Soil and Water Department, Faculty of Agriculture, Fayoum University using the standard methods described by Klute [18] and Page et al. [19] and are given in Table 1.

Table 1. Physical and chemical properties of the soil used before sowing in both seasons

| Properties               | 2014     | 2015     |
|--------------------------|----------|----------|
| Clay%                    | 29.30    | 27.50    |
| Silt %                   | 20.30    | 21.80    |
| Sand %                   | 50.40    | 50.70    |
| Texture grade            | Sandy clay loam | Sandy clay loam |
| Physical Organic matter% | 1.27     | 1.23     |
| pH                       | 7.77     | 7.79     |
| EC (dS m⁻¹)              | 7.81     | 7.82     |
| CaCO₃ %                  | 8.50     | 8.49     |
| N %                      | 0.06     | 0.06     |
| Chemical Available nutrients (mg kg⁻¹soil) |            |          |
| P                        | 19.52    | 19.20    |
| K                        | 0.37     | 0.33     |
| Fe                       | 5.66     | 5.76     |
| Zn                       | 0.79     | 0.80     |
| Mn                       | 4.95     | 5.10     |

Preparation of propolis extract (PE)

Local raw material of propolis was collected from honeybee colonies of the apiary of Faculty of Agriculture, Fayoum Governorate by scraping hives frames and entrances. Collected samples were mixed together and the active ingredients were extracted by ethyl alcohol 95% [20]. The propolis ethanol mixture was filtered and the alcohol was evaporated under vacuum (30°C) using rotary evaporator, Buchi model 011. The extract was kept cool in the refrigerator (4°C) until use. Propolis extract was diluted by water to give the final concentration required 6000, 7000, 8000 and 9000 mg/L before use. Seed treatments were carried out by soaking spinach seeds in PE or water for 12 hrs before sowing.
Table 2. Groups of substances identified in the sample of propolis, Based on Walker and Crane [21]

| Group                        | No., substance | Group                        | No., substance |
|------------------------------|----------------|------------------------------|----------------|
| - Flavonoids                 |                | Terpene and sesquiterpene    |                |
| hydroxy flavones             | 38             | alcohols and their derivatives| 7              |
| hydroxy flavanones           | 27             |                              |                |
|                              | 11             |                              |                |
| - Benzoic acid derivatives   |                | Sesquiterpene and triterpene |                |
| acids                        | 12             | hydrocarbons                 | 11             |
| esters                       | 8              |                              |                |
|                              | 4              |                              |                |
| - Benzaldehyde derivatives   | 2              | Aliphatic hydrocarbons       | 6              |
| - Cinnamyl, cinanamic acid   | 14             | Sterols and steroids         | 6              |
| and its derivatives          |                | hydrocarbons                 |                |
| - Other acids and derivatives| 8              | Minerals                     | 22             |
| - Alcohols, ketons, phenols  | 12             | Sugars                       | 7              |
| and heteroaromatic compounds |                |                              |                |
| Amino acids                  | 24             | Chalcones                    | 2              |

Seed treatment

Seeds of spinach plants (*Spinacia oleracea* L.) were obtained from Vegetable Research Institute, Ministry of Agriculture, Egypt. Seeds were sown on 1st November, for both seasons in pots (30cm in diameter and 50cm in height) each pot was filled with 20 kg calcareous saline soil. Spinach seeds were sown in each pot. Two weeks after sowing at complete germination, plants were thinned to two plants/pot.

Pot experimental

Seeds used in this study were soaking in propolis extract (PE) or water for 12 hrs, or unsoaked then grouped under 5 classes as follows:

1-Seeds soaking:

* Water (12 hrs): Seeds soaking for 12 hrs in water.
* PE 6000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 6000 mg/L.
* PE 7000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 7000 mg/L.
* PE 8000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 8000 mg/L.
* PE 9000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 9000 mg/L.

After soaking periods, seeds were air dried on filter papers overnight under the room temperature (25°C) before sowing. Each treatment contains 6 pots. The normal cultural practices for growing spinach plants were applied.

For laboratory study

Seeds used in this part of the study were treated by propolis extract (PE) and grouped under six classes as follows:

* Untreated seeds: Seeds without soaking in neither water nor PE.
* Water (12 hrs): Seeds soaking for 12 hrs in water.
* PE 6000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 6000 mg/L.
* PE 7000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 7000 mg/L.
* PE 8000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 8000 mg/L.
* PE 9000 mg/L (12hrs): Seeds soaking for 12 hrs in propolis 9000 mg/L.

After soaking period, seeds were air-dried overnight under the room temperature (25°C). The dried seeds were ground to fine powder for use in the chemical analysis of seeds.
Fertilization

All spinach plants including control were fertilized with NPK full recommended dose by the Ministry of Agriculture, Egypt. Phosphorous as triple calcium super phosphate (45-46%P$_{2}$O$_{5}$) at the rate of 75 kg/fed., (1.5g/pot) was mixed with soil before sowing. Nitrogen fertilizer was applied in the form of urea (46%N) at the rate of 100 kg/fed., (2g/pot) and 50kg/fed., of potassium sulphate (48% K$_{2}$O) (1g/pot). The amount of N and K fertilizers was divided into two equal doses, the first was added after two weeks from sowing and the second was added at two weeks later.

Measurements

Growth character

At harvest time (50-days old plants), samples of each treatment (10 plants) were taken. Plant height (cm), number of leaves/plant, fresh and dry weight of leaves/plant (g) were measured on each plant. Total leaf area (cm$^2$) of each plant was estimated by using an area meter, model Li 3000 from LI-COR, USA.

Chemical constituents

At the age of 50 days (in both seasons) samples of fresh leaves were taken for chemical determination i.e. photosynthetic pigments: chlorophyll a, b and carotenoids were extracted from fresh leaves by acetone (80%) then, their concentrations were determined (mg/ 100g fresh weight) according to [22]. Total carbohydrates mg g$^{-1}$ dry weight was determined colorimetrically according to the method described by [23]. Total and reducing sugars were determined according to [24] and recorded as mg g$^{-1}$ dry weight. Anthocyanin concentration mg/100g dry weight was determined according to the method described by [25]. Total free amino acids in fresh leaves were determined colorimetrically according to the method described by [26] and recorded as mg g$^{-1}$ dry weight. Total indoles in fresh leaves were determined colorimetrically according to the method described by [27] and recorded as mg g$^{-1}$ dry weight. Total soluble phenols in fresh leaves were determined according to [24] and recorded as mg g$^{-1}$ dry weight. Free proline concentration (mg g$^{-1}$ dry weight) was determined according to [28]. Nitrogen%, and crude protein percentage was determined according to micro Kjeldahl as described by [24] phosphorus % was determined according to [24] potassium was determined by Flame Photometer, Parkin–Elmer model 52 according to[19].

Statistical analysis

The experiment was in a complete randomized block design with 5 treatments and 6 pots as replicates for each treatment. Results were statistically analyzed using the L.S.D. a probability level of 5% for comparisons [29].

3. Result

Vegetative parameters

Effect of propolis extract (PE) on growth characters

Data presented in Tables 3, 4 indicated that propolis extract application method affected significantly growth parameters (plant height, number of leaves plant$^{-1}$, total leaf area/plant and fresh and dry weight of leaves per plant). The results showed that using seed presoaked in PE caused significant increments in growth parameters during the two studied seasons. All tested growth parameters were gradually increased by increasing propolis extract levels up to 9000 mg/L. The highest increase in yield represented in fresh weight of leaves was obtained by seeds soaking in propolis extract at 7000 mg/L, with an increase by (79.32 and 86.06%) in the first and second seasons, respectively in compared with the control (seeds soaked in water). The same trend was also observed for plant height, number of leaves/plant, total leaf area /plant and dry weight of plant leaves especially with seed presoaking propolis extract at 7000 mg/L treatment in comparison to the control plants. Applying propolis extract at 7000 mg/L effectively alleviated the adverse effects of
soil salinity on yield and its components. The highest increases were 59.78 and 74.04% for plant height, 52.70 and 60.00% for number of leaves/plant, 15.79 and 14.83% for total leaf area/plant and 79.32 and 86.06% for dry weight of plant leaves in the first and second seasons, respectively in compared with the control.

**Table 3.** Effect of propolis extract (PE) as seed presoaking agent on plant height, number of leaves/plant, total leaf area and fresh weight of leaves/plant, of spinach plants in 2014 and 2015 seasons

| Treatments            | Plant height (cm) | Number of leaves/plant | Total leaf area / plant (cm²) | Fresh weight of leaves/plant (g) |
|-----------------------|-------------------|------------------------|-------------------------------|---------------------------------|
|                       | 2014   | 2015   | 2014   | 2015   | 2014   | 2015   | 2014   | 2015   |
| Seed soaking for (12 hrs) |       |        |        |        |        |        |        |        |
| Control (water)       | 23.15  | 22.15  | 11.10  | 11.50  | 150.7  | 150.3  | 16.40  | 16.58  |
| PE 6000 mg/L          | 26.19  | 31.25  | 13.20  | 13.76  | 165.1  | 166.0  | 25.55  | 24.25  |
| PE 7000 mg/L          | 36.99  | 38.55  | 16.95  | 18.40  | 174.5  | 172.6  | 29.41  | 30.85  |
| PE 8000 mg/L          | 36.90  | 38.11  | 16.50  | 17.22  | 174.1  | 172.2  | 29.18  | 30.18  |
| PE 9000 mg/L          | 36.88  | 38.21  | 16.40  | 17.51  | 174.2  | 172.1  | 29.25  | 30.29  |
| L.S.D at 0.5%         | 4.60   | 4.40   | 2.229  | 2.72   | 6.55   | 6.33   | 4.47   | 4.28   |

**Effect of propolis extract (PE) on chemical constituents**

**Leaf pigments concentration**

Data recorded in Tables 4, 5 clearly show that the concentration of leaf pigments (chlorophyll a, b, total carotenoids and anthocyanin) was significantly increased for seed soaking with propolis extract comparing with control plants.

The data also show that seed presoaking (PE) gave the best result in chlorophyll a, b, total carotenoids and anthocyanin of spinach plants. The maximum increase was obtained with (PE) at 7000 mg/L as seed presoaking which recorded 48.04 and 49.75% for chlorophyll a, 47.62 and 59.42% for chlorophyll b, 44.64 and 44.70% for total carotenoids, 84.84 and 87.94% for anthocyanin in the first and second seasons respectively compared to the control plants.

**Table 4.** Effect of propolis extract (PE) as seed presoaking agent on dry weight of leaves/plant, chlorophyll a & b and total carotenoids concentration of spinach leaves in 2014 and 2015 seasons

| Treatments            | Leaf dry weight plant⁻¹ (g) | Chlorophyll a mg/100g F.W | Chlorophyll b mg/100g F.W | Total carotenoids mg/100g F.W |
|-----------------------|-----------------------------|---------------------------|---------------------------|-------------------------------|
|                       | 2014   | 2015   | 2014   | 2015   | 2014   | 2015   | 2014   | 2015   |
| Seed soaking for (12 hrs) |       |        |        |        |        |        |        |        |
| Control (water)       | 2.63   | 2.61   | 100.11 | 101.25 | 65.78  | 60.85  | 13.55  | 13.51  |
| PE 6000 mg/L          | 3.87   | 3.84   | 120.80 | 120.60 | 85.07  | 84.18  | 16.18  | 16.25  |
| PE 7000 mg/L          | 4.78   | 4.81   | 148.21 | 151.63 | 97.11  | 97.01  | 19.60  | 19.55  |
| PE 8000 mg/L          | 4.29   | 4.27   | 147.14 | 150.75 | 96.89  | 96.78  | 19.01  | 19.02  |
| PE 9000 mg/L          | 4.31   | 4.45   | 147.16 | 150.27 | 96.87  | 96.85  | 19.85  | 19.35  |
| L.S.D at 0.5%         | 0.63   | 0.60   | 16.02  | 16.28  | 10.54  | 10.22  | 4.25   | 4.28   |

**Total carbohydrates, total sugars and reducing sugars**

Data recorded in Table 5 clearly show that in the two successive seasons application of propolis extract at the concentrations of 6000 mg/L up to 9000 mg/L significantly increased
concentration of total carbohydrates, total sugars and reducing sugars as compared to the control plants. The best result for seed presoaking with (PE) was obtained with (PE) at 7000 mg/L which recorded 27.87 and 29.30 % increase for total carbohydrate, 25.68 and 24.08% for total sugars and 54.64 and 51.86 % for reducing sugar in the first and second seasons, respectively as compared to the control plants.

Table 5. Effect of propolis extract (PE) as seed presoaking agent on anthocyanin, total carbohydrates, total sugars and reducing sugars, of spinach leaves in 2014 and 2015 seasons

| Treatments | Anthocyanin concentration mg/100g D.W | Total carbohydrates mg/g D.W | Total sugars mg/g D.W | Reducing sugars mg/g D.W |
|------------|--------------------------------------|-------------------------------|-----------------------|-------------------------|
| Seed soaking for (12hrs) | 2014 | 2015 | 2014 | 2015 | 2014 | 2015 | 2014 | 2015 |
| Control (water) | 26.66 | 26.38 | 190.15 | 189.28 | 77.99 | 79.04 | 29.15 | 29.77 |
| PE 6000 mg/L | 37.15 | 38.21 | 216.99 | 219.55 | 89.25 | 89.21 | 32.58 | 35.74 |
| PE 7000 mg/L | 49.28 | 49.58 | 243.25 | 244.74 | 98.02 | 98.08 | 45.08 | 45.21 |
| PE 8000 mg/L | 49.02 | 49.03 | 241.15 | 243.04 | 97.50 | 97.89 | 44.78 | 44.85 |
| PE 9000 mg/L | 49.17 | 49.28 | 241.47 | 243.13 | 97.75 | 97.40 | 44.25 | 44.72 |
| L.S.D at 0.5% | 7.55 | 7.61 | 22.03 | 22.09 | 10.10 | 10.10 | 6.08 | 6.12 |

Total free amino acids, total indoles, total phenols, free proline and crude protein

Data recorded in Tables 6, 7 clearly show that in the two studied seasons, all propolis extract treatments significantly increased total free amino acids, total indoles, total phenols, free proline and crude protein in spinach leaves in comparing with the control plants. The maximum increase was obtained when 7000 mg/L of PE was used in the form of seed pre-soaking agent which resulted in 70.30 and 66.53% for total free amino acids, 79.15and 73.71 % for total indoles, 73.94 and 75.15 % increase for total phenols, 15.20 and 14.36 % for free proline and 50.00 and 48.35 % for crude protein in the first and second seasons, respectively compared with the control plants.

Table 6. Effect of propolis extract (PE) as seed presoaking agent on total free amino acids, total indoles, total phenols and free proline of spinach leaves in 2014 and 2015 seasons

| Treatments | Total free amino acid mg/g D.W | Total indoles mg/g D.W | Total phenols mg/g D.W | Free proline mg/g D.W |
|------------|-------------------------------|-----------------------|-----------------------|---------------------|
| Seed soaking for (12 hrs) | 2014 | 2015 | 2014 | 2015 | 2014 | 2015 | 2014 | 2015 |
| Control (water) | 17.21 | 17.66 | 7.58 | 7.61 | 7.79 | 7.85 | 1.71 | 1.74 |
| PE 6000 g/L | 19.98 | 20.20 | 9.99 | 9.89 | 10.22 | 10.42 | 1.80 | 1.80 |
| PE 7000 g/L | 29.31 | 29.41 | 13.58 | 13.22 | 13.55 | 13.75 | 1.97 | 1.99 |
| PE 8000 g/L | 28.15 | 28.17 | 12.96 | 12.98 | 13.21 | 13.11 | 1.90 | 1.91 |
| PE 9000 g/L | 28.25 | 28.55 | 12.96 | 12.99 | 13.08 | 13.23 | 1.92 | 1.95 |
| L.S.D at 0.5% | 2.61 | 2.59 | 2.06 | 2.08 | 2.57 | 2.69 | 0.04 | 0.03 |

Nitrogen, phosphorus and potassium concentrations

Data in both two studied seasons presented in Table 7 indicate that leaves of spinach plants contained a high concentration of nitrogen; phosphorus and potassium due to propolis extract applications condition comparing to control plants. Moreover, the chemical constituents were significantly increased with increasing propolis extract rates. The maximum increase was obtained from the application of seeds presoaking propolis extract at the rate of 7000 mg/L were 50.00 and
48.35 % for nitrogen, 48.48 and 48.48 % for phosphorous and 24.34 and 25.33 % for potassium in both seasons respectively compared to the control plants.

**Table 7.** Effect of propolis extract (PE) as seed presoaking agent on crud protein, nitrogen, phosphorous and potassium of spinach seeds in 2014 and 2015 seasons

| Treatments | Crud protein % | Nitrogen % | Phosphorous % | Potassium % |
|------------|----------------|------------|---------------|-------------|
| Control (water) | 20.62 20.68 | 3.30 3.31 | 0.33 0.33 | 1.52 1.50 |
| PE 6000 mg/L | 22.62 22.50 | 3.62 3.60 | 0.39 0.38 | 1.49 1.57 |
| PE 7000 mg/L | 30.93 30.68 | 4.95 4.91 | 0.49 0.49 | 1.89 1.88 |
| PE 8000 mg/L | 29.87 30.31 | 4.78 4.85 | 0.48 0.48 | 1.73 1.71 |
| PE 9000 mg/L | 29.93 30.37 | 4.79 4.86 | 0.47 0.48 | 1.74 1.75 |

**Chemical composition of seeds after presoaking**

**Effect of (PE) seed presoaking on seed content from total carbohydrate, total sugars and reducing sugars**

Data presented in Table 8 indicated that soaking spinach seeds in propolis extract at all the concentration significantly decreased total carbohydrate while produced a significant increase in total and reducing sugars compared to untreated seeds or soaking in water for the same soaking periods. The highest increase in total sugars was (48.56 and 50.05 %) in the first and second season respectively by PE at 7000 mg/L as compared to seeds soaking in water. The same trend was also observed for reducing sugars. The increase was (72.25 and 74.06 %) at 7000 mg/L PE soaking in the first and second season respectively as compared to soaked with seeds in water.

**Table 8.** Effect of propolis extract (PE) as seed presoaking agent on total carbohydrates, total sugars and reducing sugars of spinach seeds in 2014 and 2015 seasons

| Treatments | Total carbohydrates mg/g D.W of seeds | Total sugars mg/g D.W of seeds | Reducing sugars mg/g D.W of seeds |
|------------|-------------------------------------|-------------------------------|----------------------------------|
| Untreated seeds | 500.11 501.21 | 19.52 19.38 | 4.00 4.01 |
| Control (water) | 446.41 442.15 | 22.18 23.40 | 4.10 4.12 |
| PE 6000 mg/L | 466.21 465.11 | 24.77 24.81 | 4.52 4.60 |
| PE 7000 mg/L | 461.11 466.01 | 29.00 29.08 | 6.89 6.98 |
| PE 8000 mg/L | 460.15 461.00 | 28.11 27.99 | 6.20 6.52 |
| PE 9000 mg/L | 459.11 462.20 | 28.30 28.20 | 6.41 6.39 |

**Effect of (PE) seed presoaking on seed content from total free amino acids, total indoles and total soluble phenols**

It is clear from data showed in Table 9 that the concentration of total free amino acid, total indoles and total soluble phenols of tested seeds was increased significantly by the PE soaking at all concentration when compared to untreated seeds or soaking in water. The best result was obtained when propolis extract at 7000 mg/L was used. The highest increase was recorded 91.44 and 88.43 % for total free amino acids, 78.19 and 77.77 % for total indoles and 71.32 and 73.00 % for total soluble phenols in the first and second season respectively as compared to seeds soaking in water.

| Treatments | Total free amino acids mg/g D.W of seeds | Total indoles mg/g D.W of seeds | Total soluble phenols mg/g D.W of seeds |
|------------|----------------------------------------|-----------------------------|-----------------------------------|
| Untreated seeds | 500.11 501.21 | 19.52 19.38 | 4.00 4.01 |
| Control (water) | 446.41 442.15 | 22.18 23.40 | 4.10 4.12 |
| PE 6000 mg/L | 466.21 465.11 | 24.77 24.81 | 4.52 4.60 |
| PE 7000 mg/L | 461.11 466.01 | 29.00 29.08 | 6.89 6.98 |
| PE 8000 mg/L | 460.15 461.00 | 28.11 27.99 | 6.20 6.52 |
| PE 9000 mg/L | 459.11 462.20 | 28.30 28.20 | 6.41 6.39 |

**L.S.D at 0. 5%**

| Treatments | Total free amino acids mg/g D.W of seeds | Total indoles mg/g D.W of seeds | Total soluble phenols mg/g D.W of seeds |
|------------|----------------------------------------|-----------------------------|-----------------------------------|
| Untreated seeds | 500.11 501.21 | 19.52 19.38 | 4.00 4.01 |
| Control (water) | 446.41 442.15 | 22.18 23.40 | 4.10 4.12 |
| PE 6000 mg/L | 466.21 465.11 | 24.77 24.81 | 4.52 4.60 |
| PE 7000 mg/L | 461.11 466.01 | 29.00 29.08 | 6.89 6.98 |
| PE 8000 mg/L | 460.15 461.00 | 28.11 27.99 | 6.20 6.52 |
| PE 9000 mg/L | 459.11 462.20 | 28.30 28.20 | 6.41 6.39 |

**L.S.D at 0. 5%**

| Treatments | Total free amino acids mg/g D.W of seeds | Total indoles mg/g D.W of seeds | Total soluble phenols mg/g D.W of seeds |
|------------|----------------------------------------|-----------------------------|-----------------------------------|
| Untreated seeds | 500.11 501.21 | 19.52 19.38 | 4.00 4.01 |
| Control (water) | 446.41 442.15 | 22.18 23.40 | 4.10 4.12 |
| PE 6000 mg/L | 466.21 465.11 | 24.77 24.81 | 4.52 4.60 |
| PE 7000 mg/L | 461.11 466.01 | 29.00 29.08 | 6.89 6.98 |
| PE 8000 mg/L | 460.15 461.00 | 28.11 27.99 | 6.20 6.52 |
| PE 9000 mg/L | 459.11 462.20 | 28.30 28.20 | 6.41 6.39 |

**L.S.D at 0. 5%**
Table 9. Effect of propolis extract (PE) as seed presoaking agent on total free amino acids, total indoles and total phenols of spinach seeds in 2014 and 2015 seasons

| Treatments                  | Total free amino acid mg/g D.W of seeds | Total indoles mg/g D.W of seeds | Total phenols mg/g D.W of seeds |
|-----------------------------|----------------------------------------|---------------------------------|--------------------------------|
|                             | 2014   | 2015   | 2014   | 2015   | 2014   | 2015   |
| Untreated seeds             | 6.90   | 6.92   | 3.76   | 3.69   | 2.65   | 2.63   |
| Control (water)             | 7.12   | 7.06   | 4.40   | 4.49   | 3.09   | 3.09   |
| PE 6000 mg/L                | 9.21   | 9.33   | 4.94   | 4.84   | 4.21   | 4.35   |
| PE 7000 mg/L                | 13.21  | 13.04  | 6.70   | 6.56   | 4.54   | 4.55   |
| PE 8000 mg/L                | 12.50  | 12.66  | 5.77   | 5.74   | 4.14   | 4.27   |
| PE 9000 mg/L                | 12.64  | 12.68  | 5.76   | 5.85   | 4.18   | 4.24   |
| L.S.D at 0.5%               | 1.70   | 1.76   | 0.13   | 0.12   | 1.17   | 1.05   |

4. Discussion

It is clear with the knowledge that soil salinity reduces the various metabolic processes that are liable for traditional plant growth. The adverse result on the synthesizes of chlorophyll a, b, carotenoids, anthocyanin, sugars, total free amino acids, proline, N, crude protein and plant auxin concentration that occurred as a result of soil salt stress was reduced by treating the plants with propolis extract. In this connection, Nikolaev [30] and Salama et al. [31] reported that the increment in leaf pigments concentration of propolis extract-treated plants might well be attributed to the rise in their hormones, and/or that propolis extract enhances mineral absorption, i.e. (Fe and Mn) required for chlorophyll synthesis, since that these parts are found among mineral composition of propolis extract. [32] reported that the hyperbolic level of anthocyanin indicates an index for a decent mechanism of plant resistance towards the changes within the environmental conditions. The rise in total sugars concentration could also be attributed to the sweetening of photosynthesis by the impact of propolis extract dilated stating from its impact on seed presoaking until completely different stages of plant growth. On the opposite hand propolis extract would possibly overcome the obligatory NaCl-salt stress via accumulation of sugars. So the increase in total sugars concentration might plays a very important role in adjusting osmotic potential of the protoplasm, a conclusion that is in accordance with results obtained by [33, 15]. As well as propolis extract contain some matter like terpenoids [21, 34] from that GA3 is synthesized. The rise in total free amino acids and free proline concentrations of propolis extract-treated plants could also be appreciated for that, these plants might show higher degradation rate of proteins and/or accumulation of many amino acids ensuimg from the inhibition of their incorporation into proteins. The increment in protein concentration in propolis extract-treated plants can be attributed to that, propolis extract contains some B- group vitamins [30]. Since Tayeb [35] recorded that, vitamins of B- group acting as coenzymes, are possess some freelance roles within the biochemical processes of plants. Moreover, Rao et al., [36] proved associated increased protein synthesis with increasing in B- group vitamins accumulation might well be through functioning at the interpretation level of protein synthesis. The increment in total soluble phenols concentration in propolis extract-treated plants may well be attributed to the increment in their total sugars concentration, and/or is also thanks to the rise within the metabolic activity of those plants to synthesize shikimic acid [37]. On the opposite hand the rise in total soluble phenols synthesis in propolis extract-treated plants could indicate that, propolis extract would possibly overcome the obligatory NaCl-salt stress via accumulation of the part that constitutes with different elements (i.e. sugars, proline and total free amino acids) cellular solutes[38] for sustenance of cells state resulting in maintenance of metabolic activities in these plants. Moreover, the increment in total indoles concentration of leaves of propolis extract-treated plants is also attributed to the rise in their total free amino acids that embrace tryptophan amino acid as a precursor of IAA as shown from results of this study. The rise in macro parts in propolis extract-treated plants is also because of the presents of these parts in propolis extract. In this respect,
Walker and Crane [21] listed 149 compounds and twenty two minerals from totally different samples of propolis. Likewise as propolis extract contain many helpful mineral elements (i.e. K, Mg, Ca, Cu, Zn, Mn, and Fe) which might compensate presoaked seeds and their developed plants under the conditions of shortage of those mineral elements in carbonate soil, and/or that propolis extract could form a coat round the surface of propolis extract-presoaked seeds and make-do as a block for injurious cations and anions of (free radical) carbonate soil [34, 15]. An increasing in the measured growth characters (plant height etc.) of propolis extract-treated plants could also be attributed to the rise in indoles in these plants (Tables 5 and 6) that might induce increase in cell division and enlargement [39, 40]. Also, propolis extract contains some compounds that enhance or alter plant metabolism resulting in the rise within the leaf area [14], e.g. terpenoids which can induce the vigorous growth and/or enhance plant metabolism resulting in the rise in each fresh and dry weight [34]. Therefore, propolis extract exhibits compensatory result against the hurtful result of NaCl-salinity. This suggestion could also be because of that, propolis extract contains terpenoids [21] that have the potential to stimulate plant growth, therefore provide the plants a lucid vigor in growth to resist the adverse impact of NaCl-salinity. The positive impact of propolis extract on seed constituents of the soluble substances before planting could also be attributed to the presence of sugars among the constants of propolis extract [21] and or the rise in α–amylase activity. The rise in total free amino acids could also be due to the presence of amino acids together with tryptophan among the parts of propolis extract [21] and the, therefore, the vital increase in total free amino acids concentration could also be thanks to that, propolis extract inhibits amino acid incorporation into proteins. Furthermore, the rise in total phenols could also be attributed to the rise in metabolic activity of those tested seeds to synthesize shikimic acid [37] iatrogenic by the propolis extract and/or the rise in sugars concentration in propolis extract treated seeds. These results are typically fully in agreement with those obtained by several investigators, [7, 14-17] on different plants.

5. Conclusions

Propolis extract as seed soaking application to salt-stressed plants has been shown to reinforce plant salt stress defense responses, to act directly and/or indirectly at rising total plant performances (growth and yields) under salt stress. Thus, propolis might offer an efficient strategy to alleviate the adverse effects of salt stress through inflated N-utilization, leading to less harm to spinach growth and its protection from dangerous effects of salt stress. Therefore, propolis extract might act to alleviate the severity of salt stress on spinach plants grown on saline soils.

Conflict of Interest

The authors declare that there is no conflict of interest.

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