Impact of phosphate solubilizing bacteria on wheat
(Triticum aestivum) in the presence of pesticides

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(With 3 figures)

Abstract

Three phosphate solubilizing bacteria were isolated and identified by 16S rRNA sequencing as Pseudomonas putida, Pseudomonas sp and Pseudomonas fulva. The strains were subjected to plant biochemical testing and all the PGPR attributes were checked in the presence of pesticides (chlorpyrifos and pyriproxyfen). The phosphate solubilizing index of strain Ros2 was highest in NBRIP medium i.e 2.23 mm. All the strains showed acidic pH (ranges from 2.5-5) on both medium i.e PVK and NBRIP. Strain Ros2 was highly positive for ammonia production as well as siderophore production while strain Rad2 was positive for HCN production. The results obtained by the strains Rad1, Rad2 and Ros2 for auxin production were 33.1, 30.67 and 15.38 μg ml⁻¹, respectively. Strain Rad1 showed 16% increase in percentage germination in comparison to control in the presence of pesticide stress. Most promising results for chlorophyll content estimation were obtained in the presence of carotenoids upto 6 mgg⁻¹ without stress by both strains Rad1 and Rad2. Study suggests that especially strain Ros2 can enhance plant growth parameters in the pesticide stress.

Keywords: Pseudomonas, phosphate solubilizing bacteria, siderophore production, chlorpyrifos, pyriproxyfen.

Impacto das bactérias solubilizantes de fosfato no trigo (Triticum aestivum) na presença de pesticidas

Resumo

Três bactérias solubilizantes de fosfato foram isoladas e identificadas por seqüenciamento de rRNA 16S como Pseudomonas putida, Pseudomonas sp e Pseudomonas fulva. As estirpes foram submetidas a testes bioquímicos de plantas e todos os atributos PGPR foram verificados na presença de pesticidas (clorpirifos e piriproxifeno). O índice de solubilização de fosfato da estirpe Ros2 foi mais elevado no meio NBRIP, isto é, 2,23 mm. Todas as estirpes apresentaram um pH ácido (varia de 2,5-5) em ambos os meios, isto é PVK e NBRIP. A estirpe Ros2 foi altamente positiva para a produção de amoníaco, bem como a produção de sideróforos enquanto a estirpe Rad2 foi positiva para a produção de HCN. Os resultados obtidos pelas estirpes Rad1, Rad2 e Ros2 para a produção de auxina foram 33,1, 30,67 e 15,38 μg ml⁻¹, respectivamente. A deformação Rad1 mostrou aumento de 16% na germinação percentual em comparação com o controlo na presença de stress de pesticida. Os resultados mais promissores para a estimativa do teor de clorofila foram obtidos na presença de carotenoides até 6 mgg⁻¹ sem estresse por ambas as cepas Rad1 e Rad2. Estudo sugere que especialmente a estirpe Ros2 pode melhorar parâmetros de crescimento de plantas no estresse de pesticidas.

Palavras-chave: Pseudomonas, bactérias solubilizantes de fosfato, produção de sideróforos, clorpirifos, piriproxifeno.

1. Introduction

Plant growth promoting rhizospheric bacteria has been used for decades in horticulture and agriculture for the improvement and productivity of crops. Biodiversity pays role in the maintenance of ecosystem, it includes variety of living organisms and genetic diversity of species (Alho, 2008). Some PGPR exert advantageous effects on the plants by the process of nitrogen fixation by delivering combined nitrogen to the plant (Jetiyanon, 2015). Microorganisms perform an important role in increasing root growth, germination rate, leaf surface area water and mineral uptake percentages, crop yield and tolerance or resistance to stresses. Nitrifying and denitrifying microbes have great impact on ecosystem that it mediates nitrogen cycle (Medeiros et al., 2014). The most common bacterial genera involved in PGPR are Bacillus, Azospirillum and Burkholderia (Mangmang et al., 2015). Iron, nitrogen and phosphorous are important components for all life forms. Bacteria solubilize phosphorous which can be the substitute.
of synthetic phosphatic fertilizers (Ahemad and Kibret, 2014). Bacteria require iron for the chelation process and form siderophore complex. Phosphate solubilization and transport of ferric iron by siderophore released from PGPR increases the accessibility of different types of nutrients in the rhizosphere (Jetiyonon, 2015).

The area of soil near the root system is termed as rhizosphere (Ahemad and Kibret, 2014). Plants produce phytostimulators which increase the growth of plants, mostly cytokinins, indole-3-acetic acid (IAA), gibberellins, auxins and ACC deaminase (Abbambi et al., 2016). Some microorganisms release 1-aminocyclopropane-1-carboxylate (ACC) deaminase enzyme which produces α-ketobutyrate and ammonia from ACC. It decreases ethylene level in the plant which helps in the elongation of roots formation and improvement in seedlings survival. Indole-3-acetic acid (IAA) released from PGPR changes root growth and morphology as it facilitates plant’s nutrient uptake potentials (Jetiyonon, 2015). Usually the mechanisms used by the PGPR are for the enhancement of growth, but in stress conditions, strains mostly do not perform their functions efficiently because they have to compete in the harsh environment. However, many PGPR strains are able to tolerate the condition of stress and also have the ability to stimulate the growth of plants in stressful milieu (Parray et al., 2016). Microorganisms while living in pesticide stress, develop resistance to it. These microorganisms can effectively be used for bioremediation of pesticides contaminated sites along with their plant growth promoting attributes (Ahmad et al., 2015). Pesticides stress have been commonly applied to plants fruits, crops and vegetables for the protection all around the world. These pesticides due to persistence in the soil and also through direct exposure, harm the activities of bacteria which are able to solubilize phosphate. PSB *Pseudomonas putida* was observed for reduction in the phosphate (P) activity. Among all the pesticides, chlorpyrifos had minimum detrimental effect and can be used in the agricultural field (Kumar et al., 2015). The present study was conducted to evaluate the effect of pesticides on wheat. The pesticides chlorpyrifos and pyriproxyfen were chosen for this study as these were commonly used in Pakistan to control the pests, also it enhances crop productivity.

### 2. Materials and Methods

#### 2.1. Sample collection and bacterial isolation

Different soil samples were collected in the sterile bags from the rhizosphere of different plants. After collection, these samples of soil were processed for the isolation of bacteria able to solubilize phosphate. After making serial dilutions the samples were spread on agar plates. The isolation of bacteria able to solubilize phosphate was done on Pikovskaya’s agar (PKV) (Pikovskaya, 1948). Three phosphate solubilizing bacterial strains giving clear zones around the bacterial growth were selected for further analysis.

#### 2.2. Identification by 16S rRNA gene sequencing

The 16S rRNA gene sequencing was carried out by Macrogen, Korea. After trimming the sequences was BLAST analyzed (National Center for Biotechnology Information) in order to find the similarities between sequences. Phylogenetic trees were constructed by using MEGA 4 software by a neighbor-joining method, estimating the relationship between the halotolerant strains and reference strains.

#### 2.3. Determination of phosphate solubilization index

0.1 mL of each fresh bacterial culture was mixed in sterile distilled water and was placed on Pikovskaya, agar plates (Pikovskaya, 1948) and incubated for 7 days. By using formula, solubilization Index was measured.

#### 2.4. Phosphate estimation

PVK medium and NBRIP liquid medium was used for the estimation of the activity of bacterial strains for phosphate solubilization. 1 ml of 24 h bacterial suspension was added to each flask. All transfers of bacterial culture were carried out aseptically in triplicates and incubated at 28 °C on a shaker for 7 days. After centrifugation of samples, the supernatant was filtered and the filtrate was used for quantification of soluble Phosphate (Jackson, 1973).

#### 2.5. Effect of phosphate solubilization on pH titrable acidity

pH meter was used to determine pH change in the medium after following 7 days incubation due to the growth of phosphate solubilizing bacteria. Seven days old bacterial cultures were checked for the titrable acidity, and centrifugation of culture medium at 1000 rpm was carried for 10 minutes. Five millilitre of supernatant was titrated against 0.01N NaOH with a few drops of phenolphthalein indicator consumed per 5.0ml of culture filtrate.

#### 2.6. PGPR Attributes

##### 2.6.1. Ammonification

Ammonia production for all the isolates was checked following manual methods (Cappuccino and Sherman, 2005). Freshly grown bacterial culture was inoculated in peptone broth and incubated at 30 ± 0.1 °C for 48 h in incubator shaking was done. 0.5 ml of Nessler’s reagent was added in the test tubes after incubation. Color change was observed from faint yellow to dark brown.

##### 2.6.2. Hydrogen cyanide production

Production of hydrogen cyanide (HCN) from the bacterial isolates was observed as per methodology observed by Castric and Castric (1983). Development of light brown color was observed which indicated positive results for HCN production.
2.6.3. Siderophores production

Chrome azurol sulfonate assay agar was used for the siderophores production (Schwny and Neilands, 1987). In order to observe qualitative assay, bacterial cultures were spotted onto the blue agar and incubated at 28 °C for 24-48 h. Color change was observed for the interpretation of results, ferric ion was transferred from strong blue complex to the siderophore. Yellow- orange zones around the growth indicated positive results for siderophore production.

2.6.4. Auxin estimation

Fresh bacterial cultures were inoculated in the fifty milliliter of L-broth containing 0.1% L-tryptophan and incubated in shaker at conditions 30 ± 0.1 °C and 180 rpm for 48 h in the dark. Cultures were centrifugated at 10,000 rpm for 10 min at 4 °C. Colorimetric assay was used for the estimation of Indole-3-acetic acid (IAA) in the supernatants by adding 2 ml of Salkowski reagent in 1 ml of the supernatant. Pink color was observed and absorbance after 30 min at 535 nm in UV/Visible Spectrophotometer was read (Gordon and Weber, 1951). Regression equation was used to calculate IAA production from the standard curve and the result was expressed as µg ml⁻¹ in comparison to control.

2.6.5. ACC deaminase

Following the method of (Penrose and Glick, 2003), ACC deaminase activity for phosphate solubilizing bacteria was observed.

2.7. Effect of pesticide stress on root elongation assay

The effect of bacteria able to solubilize phosphate on growth of wheat plant was observed under pesticide stress. For this purpose watman filter paper was fitted in the petri plates. After autoclaving plates were labelled for each strain and treatment. Inoculated seeds were evenly spread in petri plates by sterile forceps. All the plates were placed in the dark for three days. Germination was observed regularly. Plates seedlings were allowed to grow in the light after germination for 10 days and parameters were recorded.

2.8. Field experiment with wheat plant

Fresh fertile soil was taken from the field of New Campus, University of the Punjab, Lahore, Pakistan. Soil was moist and brown in color. About 7-8 kg of soil was filled in each labeled pot properly. Soil was also mixed with a combination of pesticides as per recommended. Healthy seeds of Wheat (Triticum aestivum) were selected for experiment. Seeds were rinsed with tap water, discarded the floating (defective seeds) and proceeded with the healthy seeds. The seeds were dipped in 0.1% HgCl₂ solution for 5 minutes and were 3-5 times washed with sterilized water for complete elimination of HgCl₂ traces. The impact of phosphate solubilizing bacteria on wheat plant was observed under the stress condition i.e., pesticide stress. The three strains were observed with 3 treatment groups. Treatment group 1, 2 and 3. Treatment group 1 contained pesticide chlorpyrifos, treatment group 2 contained pesticide pyriproxyfen while treatment group 3 contains the combination of both pesticides (chlorpyrifos and pyriproxyfen). All these 3 treatments were compared with the control in which no pesticide was used. Seeds were sown in pots labeled with respective strains. The clean conditions were maintained during sowing. Ten seeds per pot were sown with the help of forceps. The optical density of bacterial culture for inoculation was adjusted to 0.5 at 600nm and 10 ml of this bacterial suspension was added to soil. Pots were watered equally on daily basis up-to maturation of plants. Thinning of the plants was carried out after two months and only five plants which were left to grow in each pot were selected. After thinning plant material was used for biochemical analysis.

2.9. Plant biochemical assay

2.9.1. Chlorophyll content

For the estimation of photosynthetic pigments 100 mg of fresh leaf material was homogenized with acetone (80%). The homogenate volume was kept at 10 ml and filtered to remove the plant material. The absorbance of extracted pigments was read using UV spectrophotometer at different wavelengths (470, 645 and 663nm) following Amon (1949) method.

2.9.2. Free Proline content determination

Fresh leaves of wheat plant (Triticum aestivum) were trimmed to smaller pieces. Proline concentration in leaves was determined by triturating frozen leaves of plants. Subsequently 500 mg of triturated leaf material was mixed in 5 ml of 3% sulfosalicylic acid. After centrifugation the supernatant was mixed in equal proportions (1:1:1) with glacial acetic acid and ninhydrin. After heating the reaction mixture at 100 °C in a boiling water bath for 60 min, the reaction mixture was cooled to room temperature and development of brick red colour was observed. Toluene was added to the above mixture and moved to a separating funnel and mixed thoroughly. Layer containing chromophore x was separated and its optical density at 520 nm was measured in comparison to blank. By taking 5-100 µg ml⁻¹ concentration of standard proline, proline standard curve was set.

2.9.3. Peroxidase assay

Leaf tissues (1 g) were crushed with 0.1 M phosphate buffer (4 ml). The mixture of plant was centrifuged at 4 °C for 10 min at 14000 rpm. The enzyme estimation was carried out with the supernatant. Test and control reactions were conducted side by side by mixing 0.2ml enzyme extract with phosphate buffer. Guaiacol solution was added to test reaction and left at room temperature followed by addition of H₂O₂. Blank reaction was also performed and optical density of test and control reactions was read against blank. The enzyme units were calculated as methods were described by Racusen and Foote (1965) as units per gram.
2.9.4. Acid phosphatase activity

The extraction of enzyme was carried out following methods of Iqbal and Rafique (1987). Tris HCl buffer was used to crush the plant material and supernatant obtained was used for enzyme estimation. Acid phosphatase enzyme activity was determined at pH 4.9 for one hour at 37 °C. Series of reactions were carried out for qualitative estimation including control, test, blank and standard. Optical density was measured at 510nm wavelength against water and enzyme units were calculated.

2.9.5. Protein content estimation

For the estimation of soluble protein content of plant, plant material was homogenized with phosphate buffer using pestle ad mortar and ratio of buffer to plant was 4:1. Centrifugation of samples was performed at 4 °C for 10 min at 14000 rpm. Supernatant (0.4ml) was mixed with Folin’s mixture (2ml) followed by the addition of Ciocalteu’s phenol and Folin reagent after 15 minutes at room temperature. The tubes were placed for 45 minutes at room temperature, after observing color change optical density was read at 750nm. Results were calculated with the help of standard curve.

3. Results

3.1. Isolation and 16S rRNA sequencing of Bacteria

Different samples of soil from the rhizosphere of vegetables were collected, bacterial strains able to solubilize phosphate were isolated. Three strains Rad1, Rad2, Ros2 gave promising results as compared to control for phosphate solubilization on PVK as well as NBRIP medium as shown in Table 1. The strains after isolation were directly sent for 16S rRNA sequencing using the service of Macrogen, Korea. The sequences retrieved after sequencing were BLAST analyzed and the homology was checked with the test organisms. Analysis showed that the strains belong to Pseudomonas genera (as shown in Table 2).

3.2. Phosphate solubilization potential

Phosphate solubilizing potential was measured for the isolated strains as described below.

3.2.1. Determination of phosphate solubilization index

The bacterial strains were checked for phosphate solubilization potential by observing clear zones around the bacterial colony as compared to negative control which gave no zone around the growth of bacteria. The strain Rad1 gave zone of 2.27 mm on PVK and 2.5 mm on NBRIP medium. Strain Rad2 gave good result on NBRIP medium i.e zone of 2.84 mm while on PVK medium it gave 2.42 mm. Strain Ros2 gave good result on PVK medium i.e zone of 2.6 mm was observed, while on NBRIP medium zone of 2.23 mm was observed (as shown in Table 1).

3.2.2. Phosphate estimation

The three phosphate solubilizing strains were checked for P estimation and were compared with the control. Strain Rad1 gave good results in NBRIP medium i.e 966 µg ml⁻¹ while in PVK medium 126.3 µg ml⁻¹ was observed. The most promising results were observed by strain Rad2 in NBRIP medium i.e 1163.1 µg ml⁻¹ while in PVK medium 347.4 µg ml⁻¹ was observed. Strain Ros2 gave values 955.6 and 648.3 µg ml⁻¹ in NBRIP medium and PVK medium, respectively (as shown in Table 1).

3.2.3. Effect of phosphate solubilization on pH and titrable acidity

The effect of phosphate solubilization was checked on PVK medium and also in NBRIP medium and was compared with the control. The titrable acidity value for Rad1 in PVK and NBRIP medium observed was 17.2 and 21.3, respectively. Strain Rad2 and Ros2 showed decreased titrable acidity on PVK and increased titrable acidity on NBRIP medium, respectively. Strain Ros2 gave promising results in NBRIP medium (as shown in Table 1).

Table 1. Characterization of phosphate solubilizing bacteria for solubilization potential.

| Bacterial strain | Solubilization index (µg/ml) | P estimation (µg/ml) | Titrable acidity | pH |
|------------------|-----------------------------|---------------------|------------------|----|
|                  | PVK | NBRIP | PVK | NBRIP | PVK | NBRIP | PVK | NBRIP | PVK | NBRIP |
| Control          | -   | -     | -   | -     | -   | -     | 7.0 | 7.0    |
| Rad1             | 2.27| 2.5   | 126.3| 966.0| 17.2| 21.3  | 4.4 | 5.0    |
| Rad2             | 2.42| 2.84  | 347.4| 1163.1| 8.0 | 22.5  | 4.3 | 4.9    |
| Ros2             | 2.6 | 2.23  | 648.3| 955.6| 19.6| 24.5  | 4.2 | 4.7    |

PVK: Pikovskaya agar; NBRIP: National Botanical Research Institute Phosphate.

Table 2. Identification of phosphate solubilizing bacterial strains.

| Strain code | Isolation source | Location | Identified organism       | Accession No. |
|-------------|------------------|----------|---------------------------|---------------|
| Rad1        | Raphanus sativus | Lahore   | Pseudomonas putida        | KP241947      |
| Rad2        | Raphanus sativus | Lahore   | Pseudomonas sp            | KX345931      |
| Ros2        | Rosa indica      | Lahore   | Pseudomonas fulva         | KX345930      |
3.2.4. Effect of phosphate solubilization on pH

The negative log of hydrogen ion (pH) concentration was checked for the phosphate solubilizing bacteria. pH was measured by using pH meter. Strains were compared with the control pH i.e. 7. Strains were highly acidic on both media i.e PVK and NBRIP (as shown in Table 1).

3.3. PGPR Attributes

3.3.1. Ammonia production

Phosphate solubilizing bacterial strains were checked for ammonia production i.e., development of the brown color was observed by the strains and was compared with the control in which no color change was observed after incubation. Strain Rad1 and Rad2 showed positive results for ammonia production whereas strain Ros2 showed highly positive results for ammonia production (as shown in Table 3).

3.3.2. HCN production

All the three strains were also evaluated for HCN production ability. The color change of filter paper by the strains was observed (i.e., yellow to orange). Strain Rad1 showed slight activity while strain Rad2 was highly positive for HCN production. Strain Ros2 showed negative results (as shown in Table 3).

3.3.3. Siderophore production

For siderophore production, method of Schwyn and Neilands (1987) was followed and bacteria were grown on blue agar. The strains were evaluated for siderophore production ability. Strains Rad1 and Rad2 were negative for siderophore production while strain Ros2 was highly positive for siderophore production (as shown in Table 3).

3.3.4. Auxin estimation

The strains were evaluated for auxin production in the L-tryptophan presence. Production of pink color was observed after incubation, by the strain. The results obtained by the strains Rad1, Rad2 and Ros2 were 3, 33.1, 30.67 and 15.38 µg ml⁻¹ auxin, respectively. Maximum auxin production was observed by strain Rad1 (as shown in Table 3).

3.3.5. ACC deaminase activity

The phosphate solubilizing bacterial strains were evaluated for ACC deaminase production. Strains Rad1, Rad2 and Ros2 showed ACC deaminase activity of 0.359, 0.301 and 0.278 mmol ml⁻¹, respectively. The strain Ros2 showed decreased activity of ACC deaminase. Maximum activity was observed by strain Rad1 (as shown in Table 3).

3.4. Effect of pesticide stress on root elongation assay

Maximum increase (16%) in the percentage germination under pesticide stress was observed by strain Rad1 as compared to its respective noninoculated control (see Figure 1a). With the application of strains and independent or combination of fertilizers to wheat showed significant increase in its growth as compared to noninoculated control. 88% increase in root length and 33% increase in shoot length of wheat was observed in the presence of strain Ros2 and treatment group 2 in comparison to control (see Figure 1b and 1c). There was significant 6% increase in no. of roots of wheat by strain Rad2 and treatment group 1 (see Figure 1d).

3.5. Plant biochemical tests

3.5.1. Acid phosphatase test

Strain Rad2 showed good results as compared to control with stress and without stress i.e., 0.7 and 0.5 K.A units 100ml⁻¹, respectively. Strain Rad1 showed no prominent results as compared to control. Strain Ros2 showed decreased activity i.e., 0.15 K.A units 100 ml⁻¹ with stress and 0.4 K.A units 100 ml⁻¹ without stress (see Figure 2a).

3.5.2. Chlorophyll content estimation

Reduction in chlorophyll content was observed for isolates Rad1, Rad2 and Ros2 under pesticide stress as compared to noninoculated control. Strains Rad1 and Rad2 were found to be the most effective isolates which gave values upto 6 mgg⁻¹ for carotenoids over noninoculated control (see Figure 3).

3.5.3. Peroxidase test

Peroxidase activity increased in stress conditions in case of all three strains. The most promising results were obtained by strain Rad2 i.e., 72 unit gram⁻¹ without stress and 79 unit gram⁻¹ with stress. Strain Ros2 showed results 60 unit gram⁻¹ without stress and 73 unit gram⁻¹ with stress. Strain Rad1 showed 58 unit gram⁻¹ without stress and 69 unit gram⁻¹ with stress (see Figure 2b).

3.5.3. Proline content estimation

All the strains showed decreased proline activity as compared to control. There was an increased activity of proline with stress by all the strains as compared to the treatment without stress (see Figure 2c).

Table 3. Plant growth promoting activities of phosphate solubilizing bacteria.

| Bacterial strain | Ammonia production | HCN | Siderophore production | Auxin estimation (µg/ml) | ACC deaminase (mmol/ml) |
|------------------|---------------------|-----|------------------------|--------------------------|-------------------------|
| Rad1             | +                   | +   | -                      | 33.1                     | 0.359                   |
| Rad2             | +                   | +++ | -                      | 30.67                    | 0.301                   |
| Ros2             | ++                  | -   | +++                    | 15.38                    | 0.278                   |

+ slight; ++ moderate; +++ strong.
Figure 1. Effect of phosphate solubilizing Pseudomonas inoculation and pesticide stress on germination of *Triticum aestivum* (a) percentage germination, (b) root length, (c) shoot length, (d) number of roots.

Figure 2. Effect of phosphate solubilizing *Pseudomonas* strain inoculation (a) acid phosphatase activity, (b) peroxidase activity, (c) proline concentration, (d) protein estimation of *Triticum aestivum* with and without pesticide stress.
showed production of IAA by and similar results have been reported where Rhizobia, Bacillus and Pseudomonas were able to solubilize phosphate.

According to the study done by El-Azeem et al. (2007), insoluble mineral phosphate solubilization between the range of 1.53 to 360 µg mL\(^{-1}\) and also the pH reduction from normal value of 7.1 to the range between 4.16 and 6.45 was reported. Strain Rad1 and Rad2 showed positive results for ammonia production while strain Ros2 showed strong positive results for ammonia production. Strain Rad1 showed slight activity while strain Rad2 was highly positive for HCN production. Strain Ros2 was highly positive for siderophore production. For IAA the results obtained by the strains Rad1, Rad2 and Ros2 were 33.1, 30.67 and 15.38 µg ml\(^{-1}\), respectively. Study showed low production of IAA by Pseudomonas sp. i.e., below 40.0 µg mL\(^{-1}\) (Anjum et al., 2011). The Solubilization Index (SI) was 2.7 for both of the strains of Pseudomonas (Ehsan et al., 2016). ACC deaminase activity observed by the strains Rad1, Rad2 and Ros2 showed results 0.359, 0.301 and 0.278 mmol ml\(^{-1}\), respectively. The strain Ros2 showed decreased activity of ACC deaminase. Study conducted by Ehsan et al. (2016) showed production of IAA by Pseudomonas strains in the range 54% to 88%.

In the presence of treatment group 3, strains Rad1, Rad2 and Ros2 showed percentage germination 16%, 3% and 12%. Strain Rad1 showed promising results in the presence of all the three treatment groups. When treatment group 1 was applied strain Ros2 showed 48% increase in root length while strains Rad1 and Rad2 showed no significant increase in root length. In the presence of treatment group 3, strains Rad1, Rad2 and Ros2 showed 13%, 11% and 6% increase in shoot length. The strains activity was compared with the control. Strain Rad2 showed good results for phosphatase as compared to control with stress and without stress i.e., 0.7 K.A units 100 ml\(^{-1}\) and 0.5 K.A units 100 ml\(^{-1}\), respectively. Most promising results were obtained for carotenoids production up to 6 mg g\(^{-1}\) without stress by strains Rad1 and Rad2. Under stress conditions all the strains (Rad1, Rad2 and Ros2) produced carotenoids for up to 3.5, 3.6 and 3.2 mg g\(^{-1}\) of fresh weight, respectively. Strain Ros2 showed results 60 unit gram\(^{-1}\) without stress and 73 unit gram\(^{-1}\) with stress. Strain Rad1 showed 58 unit gram\(^{-1}\) without stress and 69 unit gram\(^{-1}\) with stress for peroxidase activity. Increase activity of proline was observed with stress by all the strains as compared to without stress. Strain Rad2 gave most promising results 120 mg g\(^{-1}\) without stress and 100 mg g\(^{-1}\) with stress for protein estimation. Strain Ros2 gave results 90 mg g\(^{-1}\) without stress and 110 mg g\(^{-1}\) with stress. Use of pesticides help in causing bacteriostatic effect on microorganisms (Botelho et al., 2012). According to the study, the maximum toxicity by pesticide pyriproxyfen was observed to nodule numbers in pea, root and shoot dry of acids like gluconic acid, citric and propionic acid etc. and acidification caused by metabolic processes play important role in phosphate solubilization (Kumar et al., 2016). According to the study conducted by Rodríguez and Fraga (1999), similar results have been reported where Rhizobia, Bacillus and Pseudomonas showed slight activity while strain Rad2 was highly positive for siderophore production. For IAA the results obtained by the strains Rad1, Rad2 and Ros2 were 33.1, 30.67 and 15.38 µg ml\(^{-1}\), respectively. Study showed low production of IAA by Pseudomonas sp. i.e., below 40.0 µg mL\(^{-1}\) (Anjum et al., 2011). The Solubilization Index (SI) was 2.7 for both of the strains of Pseudomonas (Ehsan et al., 2016). ACC deaminase activity observed by the strains Rad1, Rad2 and Ros2 showed results 0.359, 0.301 and 0.278 mmol ml\(^{-1}\), respectively. The strain Ros2 showed decreased activity of ACC deaminase. Study conducted by Ehsan et al. (2016) showed production of IAA by Pseudomonas strains in the range 54% to 88%.

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biomass, shoot nitrogen and root phosphorus in greengram, leghaemoglobin, seed protein and chlorophyll content in chickpea. Root nitrogen, odule biomass root phosphorus, shoot phosphorus effects were also observed (Ahemad and Kibret, 2014).

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