Application of exotic bacterial strains as biofertilizer has always posed a constraint in the success of biofertilizer technology. Only a limited number of biofertilizers with applicability in a wide range of soil conditions is available in the market. *Pseudomonas cepacia* BAM-12 (MTCC No. 7100) (now known as *Burkholderia cepacia*), an isolate from the mungbean rhizosphere of the agricultural field of Rajasthan has shown remarkable plant growth promotion in Gujarat agricultural field soil. *B. cepacia* was found to increase the overall growth of 3 agriculturally important plants widely grown in Gujarat region, viz., mungbean, maize and rice. Increased plant growth was assessed on the basis of increased leaf number and area, biomass, chlorophyll content, profuse adventitious root branching, increased nodulation (in mungbean only), high available P in soil and overall plant growth in *B. cepacia* treated plants. Vital factor towards increased plant growth was the amount of IAA secreted by the organism (i.e. 2.65 mg IAA/100 ml of the culture filtrate) which is reasonably high in comparison to earlier reports available. Furthermore, PSB has also demonstrated bioremediation potential against harmful heavy metals such as copper, lead, nickel and arsenic by tolerating upto 25 mM indicating it’s prospective in remediating heavy metal contaminated agricultural fields.

**Keywords:** Biofertilizer; phosphate solubilizing bacteria; IAA; *Burkholderia cepacia* catecholate, mungbean, siderophore

**INTRODUCTION**

Agricultural land gets impoverished after long term cultivation, if not supplemented properly with nutrient inputs. To supplement the soil nutrient content under conventional farming systems, we need to apply high doses of agrochemicals. But during the last couple of decades chemical fertilizers have given a negative impact on the environment which in turn pollutes the ecosystem. Therefore, in order to make agriculture sustainable, it is necessary to implement a balanced and responsible use of organic agriculture (Slama et al., 2019). The principles of biofertilizers outline the similar concept where the soil health and biodiversity is built up to sustain the plant growth in longer term (Mehrzar, 2008; Jha et al., 2011). The application of environmental friendly practices is promoted by voluntary certification schemes (e.g., Global GAP or organic farming schemes) as well as by legally binding regulations (e.g., the EU Directive 2009/128) aiming at the implementation of sustainable pest management practices).

Biofertilizers help to build up the soil micro-flora and thereby the soil health. The existence of a microorganism increases the growth of plants by replacing soil nutrients (e.g., by biological N fixation, RNf) or making nutrients more available (e.g., by solubilization of phosphates) or increasing plant access to nutrients (e.g., by increasing root surface area) or releasing the plant growth regulating substances such as IAA, Gibberellins, Cytokinins etc. (VesseY, 2003). One such group of bacteria that has been extensively investigated is plant growth promoting bacteria (PGPR). PGPR are root colonizing bacteria that not only acquire benefit from the nutrients secreted by the plant root but also beneficially influence the plant in direct or indirect way like controlling phytopathogens (Parikh & Jha, 2012), resulting in stimulation of its growth (Reddy et al., 2002).

*Pseudomonads* are the most prevalent group among bacteria which dominate in the plant rhizosphere due to its high rhizosphere competence and therefore, the best colonizers (Dekkers, 1992). Several *Pseudomonads* have demonstrated their capability in promoting the overall growth of several plant species (Nezarat & Gholami, 2009; Shrivastava & Shalini, 2009; Jha et al., 2012). *Pseudomonas cepacia* promotes growth of plant mainly by solubilization of phosphate present in the soil by bringing about changes in rhizospheric soil pH and also by chelating substances which lead to solubilization of phosphates (Jha et al., 2012, 2013). In addition to phosphate solubilization, production of plant growth regulating substances of the auxin type contributes to their stimulating effects on plant growth (Dubey & Maheshwari, 2003). Intensification of energy-related and other industrial processes associated with production of wastes and by-products rich in heavy metals and other xenobiotic compounds has led to serious soil contamination of many industrial sites. The resultant accumulations of the various organic chemicals in the environment, particularly in soil, are of significant concern because of their toxicity, including their carcinogenicity, and also because of their potential to bioaccumulate in living systems (Jain et al., 2014). Thus, the intracellular concentration of heavy-metal ions has to be strongly controlled. Studies have shown that microorganisms are able to accumulate metal ions via processes such as transportation across the cell membrane (Jain et al., 2012), biosorption onto cell wall, entrapment in extracellular capsule, precipitation, oxidation-reduction reaction and biosorption to extracellular polysaccharide (Nies, 1999). Again *Pseudomonads* have received a great attention in this area also.

Although there are several marketed biofertilizers but, only limited numbers having applicability in variable soil conditions are available. In the past also there have been several reports where biofertilizer showed remarkable results in one state/country but was ineffective in another (Phospho-bacteria was successful in USSR but failed in other countries). In addition, another constraint associated with the wide application of biofertilizers is their crop specificity. Also, separate microbial formulations are required for different applications such as fertilization, bioremediation and many more. Therefore, scientists are in search of potential microorganisms exhibiting multiple properties that can be effectively applied to different crops and soil types in variable environmental conditions to promote the overall plant growth.

Realizing the needs in light of the foregoing discussion, an efficient phosphate solubilizing strain, *Burkholderia cepacia* BAM-12 (MTCC No. 7100, Genbank Accession No.1202914), an isolate from mung bean rhizosphere of the agricultural field of Rajasthan (Anamika et al., 2007) was selected for the present study. The *in vitro* and *in planta* plant growth promoting and biocontrol activities of BAM-12 were tested in 3 agriculturally important crop plants, viz., maize, mung bean and rice grown in Gujarat agricultural field soil. Further, the
heavy metal tolerance of the strain was also investigated to add more objectives to use them as bio-inoculant in heavy metal contaminated sites.

MATERIALS AND METHODS

In vitro experimentation demonstrating Plant growth promoting activity

Qualitative Phosphate solubilization assay: The ability of B. cepacia (BAM-12) to solubilize phosphate was assessed using Pikovskaya’s agar medium (containing 1% methyl red). Bacterial culture was spotted in the centre of the plate and incubated at 30ºC for 10 days. The solubilization index was calculated by measuring the zone of clearance and change of media colour from yellow to pink in the solubilization zone around the bacterial colony applying the following formula (Kumar & Narula, 1999):

\[ \text{Solubilization index} = \frac{A-B}{A} \]

\( A \) = diameter (colony + halozone)
\( B \) = colony diameter

Quantitative Phosphate solubilization assay: Quantitative determination of P solubilization by the isolate BAM-12 in broth was carried out in 100 ml of Pikovskaya’s broth (PB) medium. Composition of PB medium used is as follows (g/l of distilled water): dextrose, 10; (NH₄)₂SO₄, 0.5; MgSO₄·7H₂O, 0.1; KCl, 0.2; Yeast extract, 0.5; MnSO₄·7H₂O, 0.002; and FeSO₄·7H₂O, 0.002. Tricalcium phosphate was added at 0.1% to all the flasks separately as a source of insoluble phosphate. The media were then inoculated with 0.2 ml of respective bacterial culture (OD of 0.25 at 590 nm). Autoclaved PB medium without inoculation was considered as a control. The flasks were incubated for 5 days at 30ºC on an orbital incubator shaker at 130 rpm (Jha et al., 2012). Culture samples were harvested aseptically after every 24 hrs and water-soluble phosphate in the supernatant was determined up to 5 days according to the method of Olsen and Sommers (1982). Change in pH of culture broth was also recorded after every sampling.

IAA production by B. cepacia: Bacterial culture was tested to produce Indole Acetic Acid (IAA, an auxin) under culture conditions by growing the isolate in nutrient broth containing 0.5% glucose and 500 µg/ml tryptophan and incubated at 30ºC for 6 days. Then, 5 ml culture was taken and centrifuged at 10,000 rpm for 15 min. Supernatant (2 ml) was taken and 100 µl of 10mM O-phosphoric acid and 4 ml reagent (1 ml 0.5M FeCl₃ + 50 ml 35% HClO₄) were added to it and incubated at room temperature for 25 min. Absorbance was read at 530 nm against blank (Bric et al., 1991).

Antibiotic sensitivity test: Antibiotic sensitivity test of B. cepacia BAM-12 was performed using disc impregnated with antibiotics of different known concentrations (Hi-media, Mumbai, India). Six noted antibiotics, namely Nalidixic acid (30 mcg), Tetracycline (30 mcg), Penicillin (2 mcg), Erythromycin (15 mcg), Streptomycin (10 mcg) and Ampicillin were used. The discs were placed in circular manner over the surface of seeded BAM-12 strain in nutrient agar plates. The plates were incubated at 30ºC for 24 hrs.

Biocontrol activities

Chitinase production: Chitin agar plates were inoculated with BAM-12 and incubated at 30ºC for 3-4 days. 1% Congo red solution was poured onto the plates. The plates were incubated for 30 min at room temperature then they were washed with 1 M NaCl solution. Clear zones around the colonies on a red background were taken as the evidence for the chitinase activity.

Lipase production: Strain was tested for production of extracellular lipase enzyme by conducting the assay on tributyrin agar plates. Organism was spot inoculated on tributyrin agar plate and incubated under standard growth conditions for 4 days. A clear halo around the colony was indicating lipase production.

Siderophore production: Siderophore production by B. cepacia BAM-12 was tested using Chrome-azur II (CAS) assay medium (Schwyn & Neilands, 1987). Organism was spot inoculated on a CAS medium plate and observed for production of yellow-orange halo around it. The diameter of the halo was observed after every 24 hrs till 10 days to calculate the siderophore production efficiency.

Testing for production of Catecholate type Siderophore by Argow’s Assay: B. cepacia culture was grown in Ashby’s medium for 48 hrs. One ml of 0.5M HCl, 1 ml Nitrite molybdate reagent (prepared in sterile distilled water) was added to 1 ml of cell-free culture supernatant. Uninoculated medium served as a control. After 5 min 1N NaOH was added and observed for development of pink color whose intensity was measured at 500 nm using 2, 3-DHBA (Di-hydrobenzoic acid) as a standard (Argow, 1937).

Testing for production of Hydroxamate type Siderophore by Atkin’s Assay: Production of Hydroxamate-type of siderophores was detected and estimated in 1 ml of B. cepacia BAM-12 culture supernatant by ferric-perchlorate assay (Atkin et al., 1970).

HCN Production

Method 1(Colorimetric method): Culture was centrifuged at 7000 rpm for 15 min. at 4ºC and supernatant was passed 0.2a filter and O-phosphoric acid was added (2 ml) and was observed for development of pink color. The absorbance was read at 530 nm against blank.

Method 2 (Inverted plate technique): In this technique BAM-12 strain was grown in 10 ml of Luria-Bertani (LB) medium after incubation at 30ºC. A loopful of grown bacterial culture was streaked on the NA plate. Fungal strains were grown on potato dextrose agar (PDA) plates. An agar block of 5X5 mm² was cut from the periphery of the plate containing growing mycelium of the test fungi and was placed in the center of a fresh PDA plate. The BAM-12 streaked bacterial plate was inverted on the fungal culture plate and plates were sealed with cellophane. A control plate was also prepared which contained the test pathogen only. The plates were incubated at 30±2ºC and were observed after 3 days.

In vitro Antifungal activity of B. cepacia against Macrophomina phaseolina: ‘Antagonism’ is of common occurrence among microorganisms inhabiting the soil. Antagonistic activity of the strain BAM-12 was checked against an important devastating fungal phytopathogen viz., Macrophomina phaseolina (MTCC No. 166). Following 3 methods were employed to check the antifungal potential of B. cepacia against M. phaseolina.

Dual culture plate technique: Bacterial strain was grown in 10 ml of LB medium after incubation at 30ºC. Fungal strain was grown on PDA plates. An agar block of 0.5X0.5 cm² was cut from the periphery of the plate containing growing mycelium of the test fungi and was placed in the center of a fresh PDA plate. Different amounts (10 µl, 25 µl and 50 µl) of mid-log phase BAM-12 bacterial culture were added in the wells equidistant from the center. A control plate was also prepared which contained the test pathogen only and the wells were filled with distilled water. The plates were incubated at 30±2ºC and were observed after 7 days. Same procedure was repeated using a Minimal Essential Medium (MEM) to test the activity in deficient conditions.

Antifungal activity in broth: Fresh culture filtrate (10, 20, 30 ml) of B. cepacia was transferred separately in 250 ml conical flask containing 90, 80, 70 ml of yeast extract mannitol (YEM) broth (containing 2% sucrose) to check its antifungal potential. Actively growing mycelial discs from 5 day old culture of M. phaseoli were also transferred into each flask. The flasks were incubated at ambient temperature for 10 days at 120 rpm shaking. Flasks containing only mycelial discs in the medium served as control. After 10 days of incubation, fresh mycelial mat was harvested and dried at 85ºC for 24 hrs to constant weight for obtaining fungal dry weight. Microscopic analysis of cultures after 10 days incubation was also performed to observe the effect of bacterial culture filtrate on growth and morphology of fungal mycelium.

In vitro spor germination: Culture suspension of B. cepacia was prepared by inoculating it in LB medium and growing for 2 days to obtain sufficient cell density. Bacterial cells were pelleted to obtain the culture filtrate. One drop from culture filtrate was placed on a grease-free slide. Fungal spores were picked from 7-10 days old culture with sterilized inoculation needles and mixed with bacterial culture filtrate. The slides were placed in moist chambers made by placing two sterile filter papers each on lids and base of the petriplates. The slides with spores were then incubated at 25±2ºC for 24 hrs. Spore germination was observed under a binocular microscope after staining with cotton blue. Spores mixed in sterile distilled water only served as control. All experiments were conducted in triplicates.

In planta studies

Soil measurements

Soil testing: The experiment was conducted in a typical sandy clay loam soil and following parameters were analyzed before start of the experiment:

Soil pH, Moisture content and Available P: The method of Allen (1989) was followed to determine the soil pH using 0.01 M CaCl₂:2H₂O solution. pH meter was calibrated with standard buffers of pH 4.0, 7.0 and 9.2 prior to actual measurements.

To determine the soil moisture, the technique outlined by Piper (1950) was used. The soil moisture in percent was calculated by using following formula:

\[ \text{Moisture content (\%)} = \frac{\text{Initial weight - Final weight}}{\text{Initial weight}} \times 100 \]
To determine the available P concentration (determined before sowing and after 30 days of seed sowing), the samples were first extracted with 0.5 M NaHCO₃ (Olsen’s reagent) and then available P was determined according to Olsen & Sommers (1982).

Microbial count: The microbial count was also determined before and after 30 days of sowing. The rhizoplane and rhizosphere soil were separated by gentle tapping and composite samples were prepared. The soil samples were then air dried at room temperature and enumeration of PSB was determined by the dilution plating technique.

Bioinoculant application

Three plants, viz., rice, mung and maize were chosen for the investigation. The experiment was set in a Randomized Block Design (RBD) and 4 replicates were taken for each of the following treatments: (i) Uninoculated control and (ii) Treated: Seeds treated with the culture of *B. cepacia*.

Same treatments were taken for all the 3 plants, viz., rice, mung and maize. The experiment was carried out in open for 30 days during spring season (February-March) following normal agronomical practices described for the cultivation of three crop species. Watering of the plants was carried out on alternate days.

Preparation of inoculum: A loopful of the fresh bacterial culture was inoculated in 50 ml of PB and incubated at 30±2°C for 4 days in an incubator shaker.

Seed pelleting and sowing: 4-day old cultures having bacterial cell density of 1.6x10⁸ cfu/ml were mixed with jaggery solution separately. Mung bean, mazza and rice seeds were surface sterilized by immersing them in 0.1% of HgCl₂ solution for 3 min. The seeds were then rinsed with sterile d.w for 3 times and then blotted on a sterile filter paper, dried and pelleted in the bacterial culture. These seeds were left for air-drying. Aseptic measures were followed during the entire process and performed in a laminar air flow. Air dried seeds were sown immediately at a depth of 2 cm in 1 Kg plastic pots filled with 1 Kg sterilized soil.

To assess the plant growth, following parameters were studied:

1. Percent germination of seeds (after 5 days of sowing), Shoot and root length (after 15 and 30 days of sowing); Shoot dry weight, root dry weight, leaf area, Nodule number, chlorophyll content and available P in soil (after 30 days of sowing, i.e. at harvest). For leaf area determination, the area of each expanded leaf was calculated as:

   \[ \text{K x length x width} \]

   Where, \( K = 0.75 \) (Rugel et al., 1996).

2. Chlorophyll content: Chlorophyll content from leaf was measured by crushing the leaves in mortar pestle with 20 ml 80% acetone. The extract was centrifuged at 5000 rpm for 5 min. and supernatant was transferred to 100 ml flask. Residue was ground with 20 ml 80% acetone, centrifuged and supernatant was transferred to the same flask. The procedure was repeated until the residue became colourless. Volume was made upto 645 nm and 663 nm against the solvent (80% acetone) blank. The amount of chlorophyll present in extract was calculated in terms of mg chlorophyll per g tissue using following formula:

   1. \[ \text{mg chlorophyll a/g tissue} = 12.7(A_{663}) - 2.59(A_{645}) \times \frac{V}{1000 \times W} \]
   2. \[ \text{mg chlorophyll b/g tissue} = 22.9(A_{663}) - 4.7(A_{645}) \times \frac{V}{1000 \times W} \]
   3. \[ \text{mg total chlorophyll/g tissue} = 20.2(A_{663}) - 8.02(A_{645}) \times \frac{V}{1000 \times W} \]

   Where,
   \( A_\text{abs} \) = Absorbance at specific wavelength,
   \( V = \) Final volume of chlorophyll extract in 80% acetone,
   \( W = \) Fresh weight of tissue extracted

Testing Antifungal action of *B. cepacia* against *M. phaseolina* infestation in mung bean (in vitro studies)

Seeds were surface sterilized as previously described and then coated with 1% CMC (Carboxymethyl Cellulose) mixed with fresh bacterial (*B. cepacia*, BAM-12) culture filtrate and kept for drying for 1hr in an aseptic environment. Then seeds were wrapped with fungal mycelia and kept on a 3% water agar plate for germination at ambient temperature. Seeds treated with plain CMC and fungal mycelia served as control. Seeds were observed after every 24 hrs for symptoms of fungal infection till 8 days of treatment. Microscopic analysis of germinated seeds (seed, root, stem and leaves, wherever applicable) was also performed after every 48 hrs till 8 days to observe the effect of bacterial culture filtrate on growth and morphology of fungal mycelia.

Heavy Metal Tolerance Assay

The heavy metal tolerance capacity of *B. cepacia* was tested against 4 heavy metals, viz. arsenate, copper, lead and nickel in concentrations ranging from 0.5mm-25mM in LA plates. These metals were provided in the form of sodium arsenate, copper sulphate, lead acetate and nickel sulphate respectively. Same procedure was followed for checking the tolerance in liquid culture media (LB) also.

RESULTS AND DISCUSSION

Phosphate solubilization in Pikovskaya’s agar plate and liquid medium: The organism produced pink colored zone of clearance against the yellow background in Pikovskaya’s agar methyl red plates indicative of the acidic condition created due to secretion of organic acids by the bacteria during phosphate solubilization. The phosphate solubilization index was calculated during different days of incubation. It is inferred that the maximum increase in halo zone size occurred on the 2nd and 9th day of spot inoculation while there was regular increase in colony diameter right from the beginning till 10th day.

The appraisal of the relative efficiency of the organism in solubilizing phosphate was monitored quantitatively in Pikovskaya’s broth medium, and the results have been depicted in Fig. 1. The maximum P solubilization happened within 24 hrs of bacterial inoculation (149 ppm) that eventually led to sharp fall in pH value due to release of organic acids by the organism.

![Figure 1](image-url)
Further characterization showed that siderophore produced by the organism was of catecholate type with quite substantial concentration of 18.05 mg/L higher than other reported *Pseudomonas* spp. like *P. veronii* (isolated from the rhizosphere of *Arabidopsis thaliana*) which produced 6.4 mg/L only (Grobelał & Hilier 2017). Ferreira et al. (2019) have investigated the comparison of 5 bacterial strains producing siderophore under alkaline condition and found that *Bacillus subtilis* and *Rhizobium radiobacter* were catecholate producers. The obtained values indicate that the organism possesses high efficiency to chelate iron and eventually may be useful for acting against fungal phytopathogens.

**HCN production by B. cepacia:** HCN production by the PGPR strain is an additional advantage to its efficiency in defense during pathogen attack. In the current investigation *B. cepacia* was tested for HCN production by spectrophotometric and inverted plate methods. The organism showed pink color development on reaction with o-phosphoric acid confirming HCN production. Further, growth of the pathogenic fungi *M. phaseolina* was suppressed substantially (50%) in presence of *B. cepacia* in inverted plate method.

**Antifungal activities in plate assay and liquid culture:** Growth of *M. phaseolina* was inhibited by *B. cepacia* BAM-12 in plate assay as indicated by a clear zone around the bacterial colony (Fig. 3A). Of the 3 different amounts of bacterial culture (10 µl, 25 µl and 50 µl) tested, even 10 µl of culture was sufficient to suppress the fungal growth upto 30% after 7 days of incubation. During liquid culture studies in YEM broth, the culture filtrate of *B. cepacia* considerably inhibited the growth of *M. phaseolina* as compared to control. Development of heavy mycelial mat is quite apparent in the control flask (Fig. 3B) compared to *B. cepacia* inoculated flasks (Fig. 3C) indicative of significant fungal growth inhibition. When the quantity of culture filtrate was raised in broth, a marked decline in fungal growth was recorded as shown in Table 2, at 10% culture filtrate had inhibited 82.8% fungal growth, 20% culture filtrate inhibited 84% and 30% culture filtrate inhibited 85.35% fungal growth. The observation clearly shows that *B. cepacia* strain possesses a strong antagonistic activity against *M. phaseolina*.

**Microscopic studies demonstrating the effect of B. cepacia BAM-12 on fungal spore germination:** The bacterial strains exhibit different mechanisms and pathways to stop the aggravation of fungal pathogens. In this view, the effect of *B. cepacia* culture filtrate on spore and mycelial growth of *M. phaseolina* was checked and observed under compound microscope. The normal growth of fungal spore and septate mycelium is apparent in the control set (Fig. 4A and 4D) without bacterial culture filtrate under 40X magnification, whereas swollen and broken mycelia (Fig. 4B) in the slide containing bacterial culture filtrate can be seen. Spore initially became constricted (Fig. 4C) and eventually split and degraded in presence of bacterial culture filtrate, thus demonstrating intense antifungal potential of *B. cepacia* BAM-12.

**Table 1** Siderophore production index of *B. cepacia* during different days of incubation

<table>
<thead>
<tr>
<th>Days</th>
<th>Halo zone (A)</th>
<th>Colony Diameter (B)</th>
<th>Siderophore Production Index = A/B X 100</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.53</td>
<td>1.22</td>
<td>125</td>
</tr>
<tr>
<td>2</td>
<td>2.10</td>
<td>1.28</td>
<td>164</td>
</tr>
<tr>
<td>3</td>
<td>2.5</td>
<td>1.3</td>
<td>192</td>
</tr>
<tr>
<td>4</td>
<td>3.0</td>
<td>1.33</td>
<td>225</td>
</tr>
<tr>
<td>5</td>
<td>3.2</td>
<td>1.40</td>
<td>228</td>
</tr>
<tr>
<td>6</td>
<td>3.5</td>
<td>1.48</td>
<td>238</td>
</tr>
<tr>
<td>7</td>
<td>3.6</td>
<td>1.50</td>
<td>240</td>
</tr>
<tr>
<td>8</td>
<td>4.0</td>
<td>1.56</td>
<td>256</td>
</tr>
</tbody>
</table>

**Table 2** Antifungal activity of *B. cepacia* against *M. phaseolina* in liquid growth medium

<table>
<thead>
<tr>
<th>Culture filtrate</th>
<th>Dry Weight of <em>M. phaseolina</em> (g)</th>
<th>Control</th>
<th>Dry Weight of <em>M. phaseolina</em> (g)</th>
<th>Inoculated with <em>B. cepacia</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>10%</td>
<td>0.982±0.002</td>
<td>0.145±0.001</td>
<td>% fungal growth inhibition</td>
<td></td>
</tr>
<tr>
<td>20%</td>
<td>1.042±0.001</td>
<td>0.164±0.002</td>
<td>82</td>
<td></td>
</tr>
<tr>
<td>30%</td>
<td>1.101±0.003</td>
<td>0.172±0.002</td>
<td>84</td>
<td></td>
</tr>
</tbody>
</table>

It has been observed on several occasions that in *vitro* findings are not in coherence with the *in vivo* studies due to drastic differences in the environmental conditions (such as presence of soil and its intricate network of microflora and inorganic factors), therefore, it was quite relevant to carry out *in planta* studies in the present investigation. The strain *B. cepacia* BAM-12 has already proven its potential as a bio-inoculant in Mungbean plants in alkaline soils of Rajasthan, India (Jha et al., 2012). In order to broaden the applicability of this strain as inoculant further in terms of the environment and crop species, three agriculturally important crop plants (viz., maize, rice and mung bean) were selected for pot studies that was conducted in the soil collected from agricultural land of Anand District (Karamsad village) in Gujarat, India. During plant growth evaluation, some morphological differences like the number of leaves and shoot length were apparent among the treatments within a few days of seed sowing, and therefore, were compared only after 15 days of seed sowing whereas the additional growth parameters were assessed after 30 days of seed sowing alongwith these parameters.

**In planta Studies**

**Shoot length:** Increase in shoot length values was most striking in case of *B. cepacia* treated rice and mung bean plants, whereas the maize plants remained unaffected at 15 DAS.

**Number of leaves/plant:** PSB inoculation resulted in the increased number of leaves in rice and mung bean plants but the maize plants remained unaffected at 15 DAS.
seen in comparison to the control plants. In maize plants also notable rise (approx. 17%) in shoot length was observed in treated plants as compared to control plants. Rise in shoot length might have happened due to the combined effect of P solubilization and the production of growth-promoting substances by B. cepacia BAM-12 that eventually helped in stem expansion process (Jha et al., 2012; Chandra et al., 2018).

Plant growth parameters at harvest (30 DAS)

As evident from Fig. 5 and 6, inoculation of seeds with bacterial strain B. cepacia BAM-12 significantly improved the overall plant growth in comparison to the control in all the three crop species.

Plant height: Plant height is indicative of its growth and eventually determines the crop yield. Seed treatment with B. cepacia increased overall height of rice plants by 38.70% and of maize plants by 19.04% and significantly highest in mung bean plants by 58.75% as compared to un-inoculated control plants. The increase may be due to the production of growth promoting substances by the inoculated bacteria that carry out an important role in the stem expansion process, as discovered by Bonner (1961) and Weaver (1980). The highest growth augmentation in mung bean ascertainment greater competence of PSB strain to its native crop species than others.

Plant Fresh weight and dry weight (root and shoot): As shown in Table 3 there is a remarkable increase in fresh and dry weight of treated plants over uninoculated control. Such increase in dry weight may confer advantages to the host system with respect to its health and growth.

Leaf number and leaf area: The numbers of leaves were recorded more in the treated plants over control except in the maize plant, where the difference was not remarkable (Fig. 5). This parameter was again remarkable in the mungbean after 15 and 30 days in comparison to other plants, which again proves that it was due to the fact that mungbean is the native plant for the bacterial strain. As shown in Table 3 leaf area is denoted higher in rice and mungbean plants as compared to un-inoculated control plants (20% and 30% respectively), with only little increase in maize plants (i.e. 2%).

Table 3 Plant growth parameters of Rice, Maize and Mungbean after 30 days of sowing

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Rice</th>
<th>Treated</th>
<th>Maize</th>
<th>Treated</th>
<th>Mung bean</th>
<th>Treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plant Fresh weight (gm/plant)</td>
<td>0.092±0.002</td>
<td>0.149±0.006</td>
<td>2.17±0.004</td>
<td>2.357±0.001</td>
<td>0.679±0.008</td>
<td>1.028±0.002</td>
</tr>
<tr>
<td>Plant Dry weight (gm/plant)</td>
<td>0.033±0.001</td>
<td>0.099±0.003</td>
<td>0.467±0.005</td>
<td>0.898±0.003</td>
<td>0.286±0.002</td>
<td>0.815±0.003</td>
</tr>
<tr>
<td>Leaf Area (cm2)</td>
<td>1.63±0.007</td>
<td>1.97±0.002</td>
<td>26.24±1.20</td>
<td>26.83±1.34</td>
<td>6.63±0.05</td>
<td>8.57±0.09</td>
</tr>
</tbody>
</table>

Table 4 Chlorophyll content in Rice, Maize and Mungbean plants after 30 days of sowing

<table>
<thead>
<tr>
<th>Chlorophyll (mg/g tissue)</th>
<th>Rice</th>
<th>Treated</th>
<th>Maize</th>
<th>Treated</th>
<th>Mung bean</th>
<th>Treated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chlor a</td>
<td>1.058±0.002</td>
<td>1.505±0.003</td>
<td>0.664±0.003</td>
<td>1.026±0.003</td>
<td>1.094±0.002</td>
<td>1.095±0.001</td>
</tr>
<tr>
<td>Chlor b</td>
<td>0.612±0.001</td>
<td>0.672±0.003</td>
<td>0.381±0.002</td>
<td>0.568±0.002</td>
<td>0.432±0.003</td>
<td>0.578±0.003</td>
</tr>
<tr>
<td>Total Chl (a+b)</td>
<td>1.72±0.002</td>
<td>2.17±0.002</td>
<td>1.045±0.001</td>
<td>1.394±0.002</td>
<td>1.32±0.003</td>
<td>1.67±0.003</td>
</tr>
</tbody>
</table>

Nodule number in leguminous plant, mung bean: The nodule number in the bacteria treated mung bean plant was noted to be higher (8 in number) than that of control plants (3 in number) showing ~167% increase in nodulation.

Chlorophyll Content: The estimated chlorophyll content has been shown in Table 4. The values of total chlorophyll content were recorded again higher in treated plants as compared to control with highest increment in rice plants. Also the treated plants were optimally green in color in comparison to the untreated plants. It is possible that PSB inoculated plants may have more number of chloroplasts per cell, hence more chlorophyll, ultimately affecting the photosynthetic yield. Wang et al. (2004) found a positive relationship between chloroplast number and photosynthesis in Nicotiana sylvestris.

Deficiency symptoms in plants: In maize control plants several deficiency symptoms associated with the inadequacy of the macromolecules like nitrogen, phosphorus and potassium were observed. The symptoms were pale green colored leaves and weak stem; leaves with forward roll and scorched margins and defoliation of oldest leaves; short internodal length relatively longer leaves, marginal and tip browning of leaves. Such deficiency symptoms were insignificant in B. cepacia treated plants.

Rhizosphere bacterial Population: The introduced PSB were found to establish themselves well in the plant rhizosphere as shown by the cell count taken after 30 days of sowing (Table 5). It is evident from the results that in all the bacterial inoculated treatments, the population increased towards the maturation of the crop. The microbial population of soil was found to be higher in case of treated plants as compared to control showing occurrence of microbes in soil with inoculation of B. cepacia.

Figure 5 Growth of Rice (A), Mungbean (B) and Maize (C) after 30 days, where control plant is denoted as Cl and treated as Tr

Figure 6 Root length, shoot length and soil available phosphorus (P) concentration of rice, maize and mung bean plants after 30 DAS (days after sowing) as influenced by B. cepacia inoculation

Root branching: Root branching and root biomass significantly increased in all treated plants as compared to un-inoculated control (Fig. 5) and hence, the better developed root branching system may increase nutrient uptake by plants due to increased surface area. Indole-3-acetic acid is a phytohormone which is known to be involved in root initiation, cell division, and cell enlargement (Barazani & Friedman, 1999) and significant IAA secretion by the isolate corroborates the observed effect.
Higher number of PSB recorded after harvesting of mung bean plants at 30 days may be ascribed to the reason that bacteria took time to establish themselves in the rhizosphere and this time period varies from bacterium to bacterium. Moreover, establishment of a threshold population of viable inoculants is an important prerequisite for plant-microbe interactions like growth enhancement and biocontrol by bacteria. The results are in close conformity with those reported by Raja et al. (2002) who reported the establishment of bacterial populations in wheat inoculated with phosphobacteria.

**Available Phosphorus:** The values of available phosphorus estimated have been shown in Fig. 6 with significant higher values in treated soil as compared to control. Phosphorus is a frequently limiting macronutrient next only to nitrogen for plant growth and makes about 0.2% of plant dry weight (Schachtman et al., 1998). The increment in available phosphorus shows capability of P. cepacia to bio-ameliorate the soil P i by converting them into soluble orthophosphates that can be taken up by the plants. Highest increase was found in mung bean plants as compared to maize and rice.

**Testing Antifungal action of B. cepacia against M. phaseolina infection in mung bean (in planta studies)**

The percent seed germination was found to be 17.6% in the M. phaseolina treated control against 85% in the PSB treated, Macrophomina challenged seeds after 8 days of growth under ambient conditions (Fig. 7). Among these germinated seeds only 25% of control (M. phaseolina treated) seeds had shown little rooting compared to profuse rooting and appearance of two leaf stage in 70% mung seeds (fungi challenged and PSB treated) after 8 days of growth. The overall plant height in the control plant (without any treatment) was noted to be 15 cm while 6 cm in Macrophomina challenged control plants. On the contrary the treated seeds (with PSB culture and Macrophomina) showed 20 cm overall plant height (Fig. 9). This clearly demonstrates the growth stimulating and biocontrol potential of B. cepacia inoculation on the seeds leading to increased plant height and growth.

**Microscopic Analysis:** Microscopic analysis (400x magnifications in a compound binocular microscope) of root, stem and leaf sections of Mungbean plants revealed that the disease progression happened in pathogen challenged plants over the period of 8 days of growth. This was confirmed by the presence of disorganized, ruptured and damaged groups of cortical and endodermal cells and change in cell morphology in these sections. Blackening of the root started on the 2nd day of Macrophomina infection and advanced till the 8th day making the seed as a dead mass. On contrary the treatment of mung seeds with BAM-12 highly reduced the impact of Macrophomina attack observed as negligible blackening in roots, healthier than control plants, shoot section having intact cells without any damage.

**Heavy metal tolerance by B. cepacia BAM-12**

Agricultural soils in many parts of Asia are being contaminated by industrial wastes containing heavy metals such as arsenate, copper, nickel and lead that may have a highly deleterious effect on the microflora of soil. Thus, the efficacy of B. cepacia was tested for bioremediation of these heavy metals in different concentrations ranging from 0.5 mM to 25 mM and had shown efficiency to tolerate different concentrations for each heavy metal, i.e. 1.5 mM arsenate, 25 mM copper, 10 mM nickel and 25 mM lead respectively. Its tolerance up to 25 mM is indicative of its potential to be used in heavy metal remediation at contaminated sites, especially the agricultural fields. Studies on heavy metal tolerance by bacterial strains and its role in bioremediation (Jain et al., 2012, 2014) have revealed that the remediation of the central carbon metabolic pathway takes place in metal contaminated environments. Additionally, B. cepacia BAM-12 is also an efficient catecholate producer which might have accentuated the bioremediation process by reducing the heavy metals toxicity (Grobelak & Hiller, 2017).

**CONCLUSIONS**

In the present study a Mungbean bacterial isolate B. cepacia BAM-12 from Rajasthan agricultural field soil has demonstrated its efficacy in Gujarat agricultural field soil in terms of improving the growth of 3 highly important crops viz., Mungbean, maize and rice. Such type of crop growth improvement may be attributed to the multifaceted activities demonstrated by this isolate like direct growth enhancement (high amount of IAA production, P-solubilization), and biocontrol ability (catecholate production, chitinase and lipase activity, HCN production). Additionally the isolate has also a heavy metal tolerance trait which may be exploited for crop production in copper and lead contaminated sites. It is presumed that indigenous strains are mostly effective with indigenous plants and soil, but the isolate used in this study has demonstrated its potential in various crop species and different soil. Further, along with some other robust PGPRs it may be tested with other crop species at multilocational fields for expanding its applications.

**Conflict of interest:** The authors have no conflict of interest.

**Acknowledgements:** We are thankful to Charutar Vidyad Mandal (CVM), Vallabh Vidyanagar, Anand, India for the financial assistance and CHARUSAT University, Changa, for necessary support.

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