Enhanced Crop Productivity and Sustainability by Using Native Phosphate Solubilizing Rhizobacteria in the Agriculture of Arid Zones

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The importance of phosphate solubilizing rhizobacteria (PSB) has been well-documented as an option for enhancing sustainable agriculture. As a particular group of plant growth promoting rhizobacteria (PGPR), PSB play an important role in the soil phosphorus cycle, increasing the bioavailability for growth and plant development. This study analyses the plant growth promoting effects of 5 strains (BN0009, BN0013, BN0015, BN0024, and BN0035) out of 180 isolated from Jarava frigida (Phil.) F.Rojas (Poaceae), a native grass from the Andean Atacama desert from North of Chile. The five bacterial isolated (BN strains) were identified as non-pathogenic Erwinia sp. and show a high phosphate solubilization capacity for Ca(PO₄) ranging from 608.9 to 781.4 mg/L. Strains IAA production varies between 23.5 and 35.9 mg/L, siderophores, phosphatase (alkaline and acid) production was also observed, but none of the five isolated presented antagonism against plant pathogens Botrytis sp. and Sclerotinia sp. All isolates enhanced seed germination in Lactuca sativa and Solanum lycopersicum (excepting BN009). Additionally, all strains stimulated the early root elongation and seedling development in lettuce and tomato. Pot experiments displayed that BN0015, BN0024, and BN0035 application improved crop productivity compared to respective control. P content in plants with bacterial inoculations increased significantly compared to control in either fertilizer treatment, suggesting an improved nutrient uptake. Also, lettuce with 50% fertilization and inoculation with BN0024 equate productivity with the control 100% fertilization. Finally, we discuss these results in the context of applicability to enhance the agroecosystem productivity in arid and semiarid zones.

Keywords: PGPR - plant growth-promoting rhizobacteria, nutrient uptake, Erwinia, biostimulant, sustainable agriculture, horticulture crops
INTRODUCTION

The rhizosphere involves dynamic interactions between plant roots and soil microorganisms, where the composition of root exudates influences the nature and structure of the microbial community (Shi et al., 2011; Eisenhauer et al., 2017; López-Angulo et al., 2020). Particularly, beneficial soil microorganisms like mycorrhiza and Plant Growth Promotion Rhizobacteria (PGPR) enhance the adaptive responses and plant development by direct and indirect mechanisms (Backer et al., 2018). PGPR facilitate the bioavailability of nutrients in the soil, e.g., phosphate, considered a major macronutrient for growth and physiological plant performance (Bechtaoui et al., 2020). However, it is well-known that the bioavailable P in the soil is very limited reaching the level of 1.0 mg kg⁻¹ soil (Goldstein, 1994).

To raise assimilable P, conventional agriculture has been using chemical fertilizers. Nonetheless, the efficiency of P fertilizer throughout the world is around 10–25%, as a large proportion of the applied P is quickly transformed into insoluble forms, limiting the crop productivity and increasing environmental pollution (Kanter and Brownlie, 2019; Mallin and Cahoon, 2020).

Phosphate-solubilizing bacteria (PSB) mobilize insoluble P forms from the mineral matrix to the bulk soil where they can be absorbed by plant roots (Sashidhar and Podile, 2010). In turn, plants liberate several sources of carbohydrates, which can be metabolized for bacterial activity (Vives-Peris et al., 2020). PSB can produce various organic acids with low molecular weight, which alter the soil pH, solubilizing the phosphate from acid or alkaline soils (Kim et al., 1998; Gyaneshwar et al., 2002; Liu et al., 2020). Another possible mechanism is proton-excretion accompanying ammonium ion assimilation, which is assumed to be the most probable explanation for microbial solubilization without acid production (Illmer and Schinner, 1995). Species of the bacterial genera Aerobacter, Acetinobacter, Azospirillum, Bacillus, Burkholderia, Erwinia, Pseudomonas, and Rhizobium, have been widely reported to be able to solubilize various forms of insoluble phosphates (Ranjan et al., 2013; Goswami et al., 2016). Additionally, the use of PSB alone or in combined bioformulations with mycorrhizal fungi has been proven to be highly effective in increasing soil phosphate availability and to improve crop productivity (Singh and Kapoor, 1999; Nacoon et al., 2020). Different authors demonstrated these effects for several crops, like cotton (Qureshi et al., 2012), maize (Yazdani et al., 2009), sunflower Ekin (2010), rice (Rasul et al., 2019), or faba bean (Bechtaoui et al., 2020). Due to the activity that PSB expose by solubilizing the phosphate ion trapped in the soil constituents, the seed bacterization, and the preparation of specific biofertilizer with PSB strains are emerging as a potential strategy to promote the transition to sustainable agriculture (Babalola, 2010; Parani and Saha, 2012).

In Chile, soils of the Atacama Desert and adjacent regions are predominantly calcareous, poor in available phosphate, organic matter, and carbon content (Gómez-Silva et al., 2008). However, the presence of a high diversity of native plants adapted to arid environments and the high percentage of endemic organisms justified the consideration as one of the 35 hotspots of biodiversity around the world (Mittermeier et al., 2005). This fact suggests the presence of ecological networks that harbor bacterial strains with high biotechnical potential and beneficial effects on the productivity and growth of crops, which are grown on arid or semiarid agroecosystem. In fact, some PSB strains, isolated from arid environment, demonstrated promising applicability in agriculture (Goswami et al., 2014; Emami et al., 2020; Wahid et al., 2020).

Despite the benefits that PSB represent to the plants, the information about biotechnological applications of native microorganism, especially from Chile, still is very scarce. Therefore, the present work aims to contribute to our knowledge about PSB, their functional activity as well as potential uses in modern agriculture. We characterize PSB associated to the roots of Jarava frigida (Phil.) F.Rojas (Poaceae), a native grass that dominates the high Andean desert grasslands (Gajardo, 1994). We analyse their P solubilizing capacity and their ability to promote growth of lettuce and tomato plants; and select the best strain for a field experiment with different levels of fertilization. Finally, we discuss their applicability as biofertilizers for agriculture in arid and semi-arid zones.

MATERIALS AND METHODS

Rhizospheric Soil Sampling

For this study, we sampled a Jarava frigida population (23°02′50″S–67°39′16″W) at an altitude of 4,650 m above sea level, located in the commune of San Pedro de Atacama (Antofagasta, Chile). The sampling site is a little terrain depression with an average vegetation cover of 5%, the mean annual temperature is 7°C, and annual precipitation is ∼120–150 mm. Plants of J. frigida were harvested from the study site with a spade. Their root-sediment ball was kept intact, and they were transferred to the laboratory in sterile plastic bags. Besides the rhizoplane-root sample, soil samples were taken. Soil chemistry was analyzed by a commercial laboratory (CTSyC, University of Talca).

Soil chemistry: The pH of Atacama Desert soil samples was moderately acidic (Table 1). The total N and P level was 103 ppm v/s 0.027%, respectively. The soil samples exposed a low level of organic matter (0.63%). However, the organic N level was 0.6 times higher compared to organic P. (16 ppm v/s 10 ppm, respectively). The electric conductivity (EC), expressed as a ds/m measurement was low, but a high concentration of minerals was detected particularly for iron and copper (Table 1). In addition, a high concentration of boron was also measured, a common feature of desert regions.

Isolation of Phosphate Solubilizing Bacteria

Approximately 5 g of roots and rhizosphere (soil attached to the roots) were placed in an Erlenmeyer flask with 50 ml distilled water and shook at 120 rpm at room temperature for 2 h. Serial diluted (103–105) samples were dispersed with borosilicate glass balls on PVK plates, containing glucose 10 g/l, yeast extract 0.5 g/l, ammonium sulfate 0.5 g/l, magnesium sulfate heptahydrate 0.1 g/l, calcium phosphate 5 g/l, sodium chloride 0.2 g/l, potassium chloride 0.2 g/l, manganese sulfate 0.002 g/l,
TABLE 1 | Soil characteristics from the isolate collection site in San Pedro de Atacama (as indicated by CTSyC laboratory).

| Soil characteristics | Pasto ID 01-05-09 | Comments |
|----------------------|-------------------|----------|
| P Total              | 103.95 ppm        |          |
| N Total              | 0.027%            |          |
| P available          | 10 ppm            | Medium   |
| N available          | 16 ppm            | Low      |
| K                    | 315 ppm           | Medium−high |
| Organic matter       | 0.63%             |          |
| pH                   | 6.12              | Moderately acid |
| CE                   | 0.183 dS/m        | No risk  |

Trace minerals

- Mn: 11.02 ppm, High
- Zn: 0.75 ppm, Medium
- Cu: 2.97 ppm, High
- Fe: 9.99 ppm, High
- B: 2.65 ppm, High

Cation exchange

- Ca: 2.10 cmol(+)/kg, Low
- Mg: 0.84 cmol(+)/kg, High

ferrous sulfate 0.002 g/l, agar 15 g/l, pH adjusted to 7 (Vazquez et al., 2000). Plates were incubated for 3 days at 25°C. Afterward, solubilizing efficiency on PVK plates (triplicates) was expressed as E = halo diameter/colony diameter × 100 (Nguyen et al., 1992). All isolates with ratio > 200 were stored at −80°C.

Biochemical Characterization of PGPR Traits

Quantification of Solubilized Phosphate

The isolates were inoculated into 20 ml liquid PVK medium (as mentioned above but without agar), where Ca3(PO4)2 was replaced by KH2PO4 and incubated at 28°C for 24 h. Bacterial concentration of this liquid culture was adjusted to in 1 × 10^5 CFU in 500 µl and bacterial cells were cleaned. Finally, respectively, 50 ml of liquid PVK per isolate were inoculated and incubated for 72 h with continuous agitation (125 r/min). Every 12 h 1 ml culture was sampled aseptically for the determination of pH and soluble phosphorus. The pH was measured on a pH meter; the phosphorus availability was determined with the Mo blue method (Watanabe and Olsen, 1965). The solubilizing isolates were compared against a negative control and a not-solubilizing isolate. All measurements were taken in triplicates and the procedure was repeated 3 times independently.

Alkaline and Acid Phosphatase

To detect alkaline and acid phosphatase activity isolates were cultivated on PVK plates (composition mentioned above) for 4 days at 25°C. Afterwards, a 6 mM 4-nitrophenyl phosphate solution was added to the plates containing either acetic acid (100 mM, pH 5.4) for acid phosphatase or Tris (100 mM, pH 9) for alkaline phosphatase. In both cases, plates were incubated for 90 min., inverted and exposed to ammoniac vapor for 2–3 min. under an extractor hood. The formation of a yellow halo around bacterial colonies indicates phosphatase activity by reduction of 4-nitrophenyl phosphate (colorless) to p-nitrophenol (yellow).

Siderophores

To determine the excretion of siderophores by the strains, CAS (chrome azurol sulfur) plates were used as described by Alexander and Zuberer (1991). Bacterial strains were inoculated on the plates and incubated for 72 h. The observation of an orange halo indicates the liberation of iron from the colored Fe-CAS complex due to the presence of siderophores produced by the bacteria. Afterwards CAS-positive bacteria were grown in MM medium and incubated for 24 h (at 25°C, 120 rpm). One milliliter of the culture was centrifuged at 8,000 rpm for 5 min, the cell-free supernatant was mixed with 0.5 ml CAS solution and measured at 630 nm (spectrophotometer), as reference a 0.5 ml uninoculated MM medium and 0.5 ml CAS solution was used (Schwyn and Neilands, 1987). The percentage of siderophore units was estimated as the proportion of CAS color shifted using the formula: % Siderophore units = [(absorbance reference -absorbance sample)/absorbance reference] × 100 (Bholay et al., 2012).

IAA Production

For each strain, 10 ml liquid PY medium containing Tryptophan (0.1% p/v) were inoculated and incubated for 48 h at 25°C and 175 rpm. The production of Indole-3-acetic acid (IAA) was determined after 24 and 48 h with or without tryptophan. For each measurement, 1 ml bacterial culture was centrifuged to 0.5 ml of the supernatant 1 ml of Salkowsky reagent (1 ml FeCl3 0.5 M mixed with 50 ml 35% (p/p) HClO4; Gordon and Weber, 1951) was added and incubated in darkness for 30 min (Glickman and Dessaux, 1995). Absorbance was measured at 530 nm against a blank [PY medium + Tryptophan (0.1% p/v)]. The IAA concentration was calculated based on a standard curve calibrated with dilutions (0–100 µg/ml) of a commercial IAA reagent (Merck).

ACC-Desaminase Activity

The capacity to produce ACC deaminase was evaluated qualitatively by placing the isolates on Dworking-Foster plates with ACC as sole nitrogen source (Penrose and Glick, 2003). Bacterial growth on the plate would indicate the presence of the enzyme.

N-Fixation Ability

The capacity to fix aerial nitrogen was determined using NFb semi-solid medium according to Baldani et al. (2014).

Antagonism Capacity

To evaluate the antagonistic capacity of the strains against Sclerotina sp. and Botrytis sp., we employed the methodology of Salvatierra-Martinez et al. (2018). Briefly, PDA medium Petri dishes were used, placing in the center of each plate, a disc of fungal hyphae (7-d-old, 0.5 cm diameter) surrounded at a distance of 4 cm by five 3 ml liquid culture aliquots of each strain. The strain liquid culture was prepared in LB, grown for 24 h, and diluted 1:10. As control, plates with only a disc of Sclerotina sp.
or Botrytis sp. in the center were used. All plates were incubated at 25°C for 5 days. The inhibition of the radial growth of the fungi was calculated using the equation: 

\[ \text{ICM} = \left( \frac{C-T}{C} \right) \times 100; \]

where \( C \) = the growth area of the fungus in the control plate and \( T \) = the fungal growth area in the treatment (Fiume and Fiume, 2006). The experiment was repeated three times with 3 plates per replication for each strain.

**Strain Identity**

To determine the taxonomical identity of our strains we carried out several tests. For the Gram reaction, we used the commercial stain kit (Merck, Company) and KOH test. Additionally, photographs and observations of cell size and morphology were made for each isolate, using a light microscope and the 100 × objective lens.

Finally, we extracted genomic DNA for each pure strain, following the protocol of Yeates et al. (1998). We amplified 16S region (63F (5′-CAG GCC TAA CAT ATG CAA GTC-3′) and 1387R (5′-GGG CGG WGT GTA CAA GGC-3′; \( \approx 1,000 \) bp) with the primer pair and PCR protocol as reported by Marchesi et al. (1998). Purified fragments were sent to Macrogen for sequencing, and the resulting 16S rRNA gene sequences were compared with the NCBI database using the basic local alignment search tool (Zhang et al., 2000). The phylogenetic analysis was performed using MEGA 7 (Kumar et al., 2016). The obtained sequences are available in the NCBI database (NCBI accession numbers: MW092897, MW092898, MW092899, MW092900, and MW092901; not released yet).

**Promotion of Plant Growth**

The effects of the bacterial isolates on plant growth were evaluated using Lactuca sativa L. (lettuce) and Solanum lycopersicum L. (tomato).

**Seed Germination Assays**

Seeds [L. sativa (lettuce) and S. lycopersicum (tomato)] were sterilized with sodium hypochlorite and vernalized for 48 h at 4°C. Afterwards, they were submerged in liquid bacterial culture [LB medium, \( \text{OD} = 0.8 \approx 10^7 \text{ CFU} \)] for 5 h at room temperature on a shaker (180 pm). Twenty-five prepared lettuce or 15 prepared tomato seeds were displaced in a Petri dish on humid filter paper (each isolate in quadruplicate, negative control without bacteria). Petri dishes were incubated horizontally at 20°C for 3.5 days (lettuce) and 14 days (tomato), respectively. Observations on germination progress were made every 4 h for lettuce and every 8 h for tomato. Root length was determined at the end of the respective incubation period, using the scale: no root, radicle emerged, radicle <1 cm and root > 1 cm.

**Pot Experiment With Lettuce**

For this experiment, the five selected bacterial strains (BN0009, BN0013, BN0015, BN0024, and BN0035) were grown on nutrient agar initially. A single colony was transferred into 250 ml flasks containing nutrient broth and grown aerobically on a shaker (150 pm) for 24 h at 28°C. The bacterial suspension was then diluted in sterile distilled water to a final concentration of \( 10^8 \) CFU/ml. Sterile seeds were submerged for 5 h in this bacterial solution. Inoculated seeds were placed in a pot with a 1:1 sterile mixture of agricultural soil and peat. For each isolate, 10 pots were prepared (replicates) and located randomly. Pots were maintained in a climate chamber at 20°C for 28 days (photoperiod: 16 h light, 8 h darkness). The pots were irrigated daily 5 ml with distilled water. As negative control, 20 pots containing seeds without bacteria were cultured under the same conditions. Photographs were taken every 5 days. At the end of the incubation period, root and leaf area (using WinRhizo), number of leaves, as well as root and leaf fresh and dry weight were recorded. Root and shoot portions of lettuce plants were dried separately for 5 days at 65°C (constant weight).

**Lettuce Field Experiment**

Based on the previous results, one strain was selected for the lettuce field experiment. Just before transplantation, lettuce seedlings (28 days old) were soaked for 1 h in a bacterial suspension. Therefore, the strain was grown in liquid LB medium in a flask on a shaker (150 pm) for 24 h at 28°C and this bacterial culture was diluted afterwards to the final concentration of \( 10^8 \) CFU/ml. The strain was re-inoculated 4 and 8 weeks after the transplant, via drip irrigation adjusting the concentration at the dripper to \( 10^8 \) CFU/ml in 10 min (a separate irrigation system was installed for these applications). We considered two fertilization treatments with respect to the literature indication for crop management of lettuce (INIA, 2012): 100% (NPK ratio 100-100-80) and 50% (NPK ratio 50-50-40), using KNO3, \( \text{NH}_4\text{H}_2\text{PO}_4 \), and \( \text{NH}_4\text{NO}_3 \) diluted in water for applications during drip irrigation. In total, we established four treatments: control 100% fertilization, control 50% fertilization, BN0024 100% fertilization and BN0024 50% fertilization. After 3 months lettuce was cropped and 10 individuals per treatment (randomly picked) were used for analyses. Growth parameters (root and aerial weight, root area) and biochemical parameters (leaf P content (Hylander et al., 1996), protein content (Bradford, 1976), and chlorophyll content (Porra et al., 1989) were measured.

**Statistical Analyses**

The experimental data were analyzed regarding homogeneity (Pettitt test), variances (Barlett test), descriptive statistics (standard variation and error, median, mean, asymmetric coefficient, range, maximum, and minimum value). Afterwards, Tukey’s range test and ANOVA were applied to determine means (from the bacterial treatments) that are significantly different from the control and/or other treatments \( (P < 0.05) \). As statistical and graphical software, XLSTAT, and SigmaPlot were used.

**RESULTS**

**Screening for Phosphate Solubilizing Bacteria and Quantification of Solubilized Phosphate**

From Jarava frigida rhizosphere samples, 180 isolates were obtained on PVK plates. The isolates showed different colony morphology features such as color, density and colony elevation (Supplementary Table 1). A qualitative determination of the
phosphate solubilization capacity was carried with the whole collection using PVK agar plates. Measurements of solubilization efficiency E ranged between 110 and 300, with an average of 172. Twenty-seven isolates demonstrated a solubilization efficiency of over 200 and were further characterized regarding their capacity to solubilize P in liquid medium (Supplementary Table 1). The five most efficient solubilizing isolates were selected for further analyses due to their outstanding E-values of 260 to 300 (Figure 1A). These isolates were labeled as BN0009, BN0013, BN0015, BN0024, and BN0035. Their colonies were described as round and creamy in color (Figure 1B). All BN isolates exposed a high growth capacity on LB medium, with a density peak after 16 h of incubation (Figure 1C). At the same time (16 h), the growth of these five isolates in PVK liquid medium containing a P soluble source is noticeably reduced (Figure 1D). For these selected isolates a quantification of solubilized P in a liquid medium was carried out, which demonstrated different levels in their capacity dissolving insoluble phosphate (Figure 2). The highest solubilization activity was observed between 36 and 48 h when 608.9–781.4 mg/L P were liberated from BN strains. A longer incubation time did not further increase the solubilized phosphate by each isolated. On the contrary, by the end of the experiment (72 h), the concentration of soluble P decreased slightly in all strains. At the same time pH descends from the initial 7.0 to values ranging from 4.3 to 4.0, apparently in negative correlation to the amount of dissolved P. BN0024 and BN0035 solubilized 781.4 and 746.5 mg/L, respectively, the highest amount of P as bacterial culture (Figure 2). Nonetheless, when the capacity of single cells to solubilize phosphate was calculated, the most efficient strain is BN0015 with 10 pg/CFU after 36 h of growth (Figure 3).
FIGURE 2 | Kinetics of phosphate solubilization and pH changes of the five BN isolates in time, measurements every 12h during 3 days of incubation.

Evaluation of PGPR Traits
With respect to the taxonomy and identity, all five BN strains were determined as Gram-negative. Based on the 16S sequences, the strains belong to the genus Erwinia sp, with the highest identity scores (> 98.5%) for Erwinia pyrifoliae strain 8557, Erwinia billingiae strain HMF7415 and strain P3_3, a bacterial endosymbiont of Curculio koreanus (Supplementary Figure 1). Our strains are closely related to one another but are not clones. The phylogenetic analyses did not allow us to determine the species.

Furthermore, several PGPR characteristics were analyzed (Table 2). First, the ability to produce IAA, an important
phytohormone for plant growth promotion was assessed and detected for all strains. The IAA production with tryptophan enriched medium ranged from 23 to 54 mg/L (Table 2). Among all BN isolates, BN0009 and BN0035 produced the highest IAA amounts (31 and 54 mg/L, respectively). BN0015, the most efficient P-solubilizer was found to produce an average IAA value (23.5 mg/L). Without tryptophan the IAA production was noticeable lower in all strains, reaching concentrations between 1.4 and 10.3 mg/L (Table 2).

All isolated strains demonstrated activity of alkaline and acid phosphatases and were capable of nitrogen fixation and siderophores production, in the latter values ranged between 0.5 and 10% siderophore units (Table 2). Additionally, only BN0009 and BN0013 were able to grow on ACC medium suggesting that both isolates possess ACC Deaminase activity (Table 2).

None of the five strains presented antagonism against Sclerotina spp and Botrytis spp. (Table 2).

### Seed Germination Assays

Seeds of *L. sativa* (lettuce) and *S. lycopersicum* (tomato) seeds were inoculated with the selected strains to evaluate their effect on germination. All BN isolates accelerated the germination of *L. sativa* seed compared to the non-treated control, but the rate of enhancement varied among the bacterial strains (Figure 4A).

Nonetheless, BN0013 accelerated the germination most (4.6x), reaching up to 93% after 24 h of incubation vs. the control with 20% at the same time (Figure 4A). Additionally, early root development and extension were analyzed 32 h after inoculation (Figure 4B). In *L. sativa* seeds, all isolates fostered early root development compared to control. In seeds treated with BN0009 and BN0013 a 78% (51% of seed with radicle <1 cm, 27% with root >1 cm) and 88% (45% of seed with radicle <1 cm, 43% with root >1 cm), respectively, demonstrated an enhanced root development compared to the non-treated control seeds (14% of seed with radicle <1 cm, 0% root >1 cm) (Figure 4B).

In the germination assay with *S. lycopersicum* seeds, again all isolates accelerated the germination in 2.5 and 2 times in 6 days with respect to the non-treated control (Figure 4C). Meanwhile, BN0024 anticipated germination, reaching the first peak after only 4 days of incubation (45% of the seeds germinated), BN0009 significantly increased the germination rate compared to the non-treated control. After 7 days of incubation, in the non-treated control 50% of the seeds were germinated, whereas the germination rate in inoculated seeds varied between 72 and 92% (Figure 4D). Similar to lettuce, all BN isolates enhanced early root development in *S. lycopersicum* seeds. 45% (BN0024) to 60% (BN0009) of seeds with bacterial treatments already presented roots >1 cm, compared to the control with only 8% of the seeds (Figure 4D).

Finally, the root length of lettuce and tomato seedlings was measured after the respective incubation period of 3.5 and 14 days, respectively (Figure 5). In both crops, the BN isolates induced a significant root elongation compared to the non-treated control. For lettuce, BN0013 caused the longest root extension with 2.3 times compared with the control (Figures 5A,B), followed by BN009 and BN0015. In tomato, the bacterial effect is more equal. However, isolates BN0013 and BN0024 rise with an average root length of 2.0 and 2.2 times compared to the control (Figures 5C,D).

### Pot Experiment With Lettuce

Further insights about growth promotion beyond germination produced by the five strains were evaluated in a lettuce pot experiment. Inoculated seeds were planted and maintained for 35 days under controlled conditions. At harvest, root area, leaf area, number of leaves, fresh, and dry weight were measured. All isolates significantly increased root area (1.4–1.8x), leaf area (1.5–1.7x), and the number of developed leaves (1–2 leaves).
FIGURE 4 | Characterization of the bacterial effect on germination and early root growth in lettuce and tomato; (A) germination over time in lettuce; (B) comparison of bacterial treatments regarding root growth categories in lettuce seedlings at 32 h of incubation; (C) germination over time in tomato; (D) comparison of bacterial treatments regarding root growth categories in tomato seedlings at 7 days of incubation.

compared to the control (Figure 6A, Table 3). Also, the effects were similar when comparing the strains. Seeds treated with BN0024 developed on average one leave more than control, which implies an advance in seedling development of ~1 week. The results for fresh and dry weight after inoculation with BN isolates were more differential (Figure 6B, Table 3). All isolates presented a significantly higher root fresh weight (1.9–2.3x), but the root dry weight was similar to the control. Three strains (BN0015, BN0024, and BN0035) stimulated significantly the leaf development in both fresh and dry weight compared with the control (1.8, and 2x, respectively) (Figure 6B, Table 3). Overall, the inoculation with BN0015, BN0024, and BN0035 produced the strongest promotion of plant growth and development on Lactuca sativa during the pot experiment.

**Lettuce Field Experiment**

Based on the previous data, BN0024 was selected for a lettuce field experiment with root inoculation during transplant from nursery to field and two fertilization levels. The lettuce plants responded to both factors (inoculation/fertilization) of the field experiment (Figure 7, Table 4). The effect of bacterial inoculation on plant growth was higher at 50% fertilization, where all evaluated parameters increased significantly (Table 4). With 100% fertilization the evaluated parameters also improved in plants with application of BN0024, nonetheless only root area, P content and protein content were significantly higher compared to control. The P content in plants with bacterial inoculations was significantly elevated compared to control in either fertilization treatment.

Finally, the application of the strain BN0024 with 50% fertilization reached a crop productivity and nutrient content similar to the control with 100% fertilization (Table 4).

**DISCUSSION**

In this work, rhizosphere soil samples from Jarava frigida (Phil.) F.Rojas (Poaceae) were screened for phosphate solubilizing rhizobacteria, which exhibit additional PGPR traits. The
rhizospheric soil showed a low content of organic matter (0.63%), lower electric conductivity and high content of iron and copper, as described for Andean desert soils (Mandakovic et al., 2018).

Among the 180 phosphate solubilizing isolates, five efficient BN isolates were selected for further studies. Each of these five isolates demonstrated a high P-solubilization capacity, ability to N fixation, as well as the production of siderophores and IAA, which are used as indicators for a positive growth promotion aptitude (Backer et al., 2018; Wahid et al., 2020). However, but none of them showed antagonism response against fungal pathogens. The five strains, denominated as BN0009, BN0013, BN0015, BN0024, and BN0035 were identified as Erwinia sp., and did not demonstrate pathogenic characteristics. In fact, non-pathogenic Erwinia sp. strains were previously described as phosphate solubilizes by different authors, like Goswami et al. (2016) and Sagar et al. (2018).

In liquid medium each of five BN strains, induced an inverse relationship between pH and the solubilized P concentration, suggesting that the acidification facilitates the inorganic phosphate solubilization (Khan et al., 2007). This capacity is probably complemented with the enzymatic activity of alkaline and acid phosphatases that use organic phosphate as a substrate to convert it into inorganic form (Walpola and Yoon, 2012; Zhao et al., 2013). Additionally, the increased plant elongation (plant height and root length) can be associated with higher nutrient absorption, particularly phosphorous, which can induce cell elongation and multiplication Richardson and Simpson (2011).

Another important feature in the rhizobacteria—plant interaction, is their ability to produce phytohormones. We
looked especially at IAA, which is involved in the formation of lateral roots and root hair, enhancing the nutrient absorption from soil (Qin and Huang, 2018). At least four pathways have been described for the biosynthesis of IAA in PGPR strains, being one of them tryptophan Trp-independent and two of them (Trp)-dependent (Gusain and Bhandari, 2019). Our five BN isolates produced between 23 and 54 mg IAA/L under the presence of tryptophan, concentrations which could cause important effects on the growth of the host plant. Actually, up to 80% of IAA-producing bacteria colonize root surfaces and stimulate the root system as a response to higher IAA concentrations (plant exogenous and endogenous synthesis) (Patten and Glick, 2002). The five isolated strains also were able to produce IAA in absence of tryptophan, which suggests the presence of a tryptophan-independent pathway (Ribeiro and Cardoso, 2012).

All selected isolates demonstrated stimulation on lettuce and tomato germination (Figure 4). All strains accelerated the germination significantly compared to the control and generated a fast root extension in their early growth phase. These results coincide with other PSB studies, using strains alone or in combination, where a strong stimulation in radicle emergence from the seed is observed in plants with economical interest, e.g., rice, asparagus bean, and mung bean (Duarah et al., 2011; Arun et al., 2012).

In the lettuce greenhouse experiments, all five strains enhanced significantly the plant growth regarding root and leaf area, fresh, and dry weight as well as leaf numbers. These results are concordant with other authors who studied the effects of PSB on different crops: canola (Pandey et al., 2006), peanuts (Goswami et al. (2014), cotton (Qureschi et al., 2014), maize (Yazdani et al., 2009; Shirinbayan et al., 2019), and rice (Rasul et al., 2019).

Despite the variety of in vitro and pot experiments with PSB and PGPR, few studies are available, where their effects were evaluated under productive conditions in field. Here we challenged Erwinia sp. strain BN0024, selected due to its constant growth promotion results, in a field experiment with lettuce, comparing two fertilization treatments (50 and...
100% of the recommended crop fertilization). The application of BN0024 improved crop productivity in both fertilization treatments. Particularly, the P content in plants with bacterial inoculations was significantly elevated compared to control in either fertilization treatment, suggesting an improved nutrient uptake (Rezakhani et al., 2019; Bechtaoui et al., 2020). In addition, lettuce with 50% fertilization and inoculation with BN0024 equate the productivity of the control plants with 100% fertilization. Recently, similar experiences reducing the phosphate fertilizer requirements of crops between 50 and 75% have been reported for other soils, e.g., Brazilian Latosol with sugarcane (Saccharum sp.) production integrating PSB (Rosa et al., 2020), Chinese Troporthents and Rhodustuts with tea (Camellia sinensis (L.) O. Kuntze) production integrating PGPR (Tennakoon et al., 2019). Our results gain special importance in the context of P availability in agricultural soil of semiarid and arid zones, which could be mediated integrating PSB application in the crop management (Bechtaoui et al., 2020; Wahid et al., 2020) reducing environmental pollution (Mallin and Cahoon, 2020) and production costs for agriculture.

Beyond aridity, soils of arid zones possess other complex characteristics (pH, low soil organic matter, low availability of macronutrients, bound by soil chemistry), which need to be addressed in agriculture management. While aridity is compensated by the use of irrigation systems, the low availability of macronutrients like phosphorus, quickly bound to the soil matrix and not accessible for plant growth, still is a major issue in agriculture and the use of P solubilizing PGPR can offer an alternative to improve fertilization efficiency—as demonstrated by our results. Our results encourage the use of PGPR for these
purposes, improving not only the efficiency of P-fertilization, but also allowing to improve crop quality and productivity as well as reduce costs.

Future studies will focus on two aspects: (1) further biochemical, physiological, and molecular characterization of the five *Erwinia* sp. isolates to understand more entirely their metabolism and interaction with plants; (2) development of application strategies for these strains as part of integrated crop management. This information will enable the design of eco-friendly alternatives to stimulate the productivity of agriculture carried out under the limiting conditions of semi-arid and arid zones.

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: NCBI accession numbers: MW092897, MW092898, MW092899, MW092900, and MW092901.

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**AUTHOR CONTRIBUTIONS**

AS, JB, and VO designed the research. SM, AR, BA, and PM sampled and performed laboratory work under the guidance of AS and CJ. JA, MG, and CS carried out data analysis. All authors wrote the manuscript, designed Tables and Figures, revised the manuscript, and approved the final version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fsufs.2020.607355/full#supplementary-material
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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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