Identification and Characterization of the Phosphate-Solubilizing Bacterium *Pantoea* sp. S32 in Reclamation Soil in Shanxi, China

Qian Chen and Shanjiang Liu*

Institute of Plant Nutrition and Resources, Beijing Academy of Agriculture and Forestry Sciences, Beijing, China

Phosphate solubilizing bacteria (PSB) can convert insoluble forms of phosphorus (P) to accessible forms. Five highly efficient PSB strains, H22, Y11, Y14, Y34, and S32, were screened and isolated from an alfalfa rhizosphere in heavy metal-contaminated reclamation area in Shanxi Province, China. Based on morphological observations, 16S rRNA sequencing, cellular fatty acid composition analysis, and the BIOLOG test, H22, Y11, and Y34 were identified as *Pseudomonas* sp., while Y14 and S32 were identified as *Pantoea* sp. Among them, S32 showed the highest P-solubilizing efficiency in culture medium containing Ca$_3$(PO$_4$)$_2$, lecithin, and powered phosphate rock. The culture medium conditions to obtain the highest P-solubilization efficiency were optimized as follows: the culture temperature was 30°C; the incubation time was 5 days; the initial pH was 7.0; and glucose served as the carbon source. Furthermore, the P-solubilization efficiency of S32 in media containing CaHPO$_4$, lecithin, phosphate rock (PR), FePO$_4$, or AlPO$_4$ was determined to be 18.38, 3.07, 0.16, 0.51, or 2.62%, respectively. In addition, the acid and alkali phosphatase activities of S32 were tested as 6.94 U/100 mL and 4.12 U/100 mL, respectively. The soil inoculation experiment indicated that inoculation with S32 resulted in an obvious improvement in the available P of both the experimental and reclaimed soil. The rice seedling growth experiment also suggested that the application of S32 significantly increased the plant height, biomass, root growth, and P uptake of rice in both experimental and reclaimed soil. Taken together, the isolated S32 strain showed high P-solubilization capacity for both Pi and Po, and its ameliorative effect on reclaimed soil recovery provides the theoretical basis for crop development in the reclaimed soil of mine field.

Keywords: PSB, *Pantoea* sp., phosphatase activity, reclaimed soil, pot experiment

INTRODUCTION

Phosphorus (P) is the second most important nutrient for plant growth, accounting for 0.2% (w/w) of plant dry weight (Maharajan et al., 2018). P plays an irreplaceable role in the ecosystem by participating in most aspects of energy metabolism, nucleic acid and protein synthesis, and kinase regulation (Nesme et al., 2018). The average P content in soil is nearly 0.05% (w/w) with the main two forms being inorganic P (Pi) and organic P (Po). Nevertheless, only 0.1% of P can be utilized by plants, rendering available P a restrictive factor for plant growth (Lambers and Plaxton, 2018). Phosphate anions in chemical fertilizer available to plants are extremely reactive and become...
fixed through interactions with Ca$^{2+}$, Fe$^{3+}$, and Al$^{3+}$ ions in the soil to form insoluble phosphate salt complexes; however, the plant utilization efficiency for P in chemical fertilizers is only 5–25% (Schnug and Haneklaus, 2016), leading to P enrichment in the soil and the loss of soil fertility. These realities make increasing the P utilization rate for plant growth an urgently needing situation.

Phosphate solubilizing bacteria (PSBs) convert unavailable P (both Pi and Po) into available P to satisfy the requirements of plants through dissolution and absorption. According to the various P-dissolving patterns, PSBs can be divided into two classes: (1) Pi-solubilizing microorganisms that secrete organic acid to dissolve Pi compounds and (2) Po-mineralizing microorganisms that secrete phosphatase to enzymatically mineralize Po compounds. The application of both classes of PSBs in soil decreases the pH of the soil and forms a P-offering microarea around the plant rhizosphere, consequently improving the P supply available to the plant and strengthening the activity of other beneficial microorganisms, such as *Rhizobium* and *Trichoderma*. These applications promote the absorption of nutritive element ions.

Recently, following the development of energy bases and mines explored in Shanxi Province, China, a vast amount of cultivated land was contaminated with gangue, mineral waste, heavy metal, and rock debris. Approximately 68 km$^2$ of cultivated land lost their crop productivity. Therefore, rational and scientific measures for soil reclamation in the mining area should be developed and applied without delay. In this study, to acquire a high PSB strain and improve the soluble P available to plants in the reclaimed soil, five PSBs were screened from the alfalfa rhizosphere soil of the reclaimed area in the mining area of Xiaoyi town in Shanxi Province, China. Their P-solubilizing capacities under different conditions were investigated. The effect of PSB S32 on P solubility in the reclaimed soil recovery, plant and root growth, and the P uptake of rice were also evaluated. The present data suggest that the application of the isolated PSB would be of great importance in the bioremediation of reclaimed soil in mining areas.

### Physical and Chemical Properties of Reclaimed Soil

The reclaimed soil was aseptically separated from the roots to assess the physical and chemical properties. The pH was measured in a 2:1 watersoil suspension with a pH meter (Corey, 1971). Soluble P was extracted by the bicarbonate method and was then analyzed by the Molybdenum blue method (Colwell, 1963). The organic matter content was measured using the potassium dichromate colorimetric method (Nelson et al., 1996). The available potassium content was determined using the flame photometer method (Gammon, 1951). The content of nitrate (NO$_3^-$) and ammonia nitrogen (NH$_4^+$) extracted by a potassium chloride solution was determined with an AutoAnalyzer 3 (AA3) equipped with NH$_4^+$ and NO$_3^-$ channels (SEAL, Germany) (Crooke and Simpson, 2010). Briefly, 1–2 g of soil that had passed a 2-mm sieve were placed into a 50 mL Erlenmeyer flask. Twenty mL of KCl solution was added to the flask. Samples were shaken on a horizontal shaker for 1 h and filtered through Whatman No. 2 filter paper. Filtrates were analyzed for NH$_4^+$ and NO$_3^-$ by the Molybdenum blue method (Colwell, 1971). Soluble P was extracted by the bicarbonate method and filtered through Whatman No. 2 filter paper. Filtrates were then analyzed by the Molybdenum blue method (Colwell, 1971). Extractions of all soils were repeated three times. Data are presented as the means ± standard deviation. The physicochemical properties of the soil samples are shown in Table 1.

### Culture Medium

**Pi culture medium** (Sahu and Jana, 2000) consisted of 10.0 g glucose, 0.5 g (NH$_4$)$_2$SO$_4$, 0.5 g yeast extract powder, 0.3 g NaCl, 0.3 g KCl, 0.03 g FeSO$_4$·7H$_2$O, 0.3 g MgSO$_4$·7H$_2$O, 0.03 g MnSO$_4$·4H$_2$O, 5.0 g Ca$_3$(PO$_4$)$_2$, 1000 mL distilled water, and 20.0 g agar, pH 7.0–7.5.

**Po culture medium** (Surange et al., 1997) consisted of 10.0 g glucose, 0.5 g (NH$_4$)$_2$SO$_4$, 0.5 g yeast extract powder, 0.3 g NaCl, 0.3 g KCl, 0.03 g FeSO$_4$·7H$_2$O, 0.3 g MgSO$_4$·7H$_2$O, 0.03 g MnSO$_4$·4H$_2$O, 1.0 g CaCO$_3$, 0.2 g lecithin, 1000 mL distilled water, and 20.0 g agar, pH 7.0–7.5.

**Nutrient agar culture medium** (Deshmukh, 2007) consisted of 10.0 g peptone, 3.0 g beef extract, 5.0 g NaCl, 1000 mL distilled water, and 15–20 g agar, pH 7.2–7.4.

**Trropic soy broth (TSB) culture medium** (Mishra and Goel, 1999) contained 15.0 g tryptone, 5.0 g soy peptone, 5.0 g NaCl, and 1000 mL distilled water, pH 7.2–7.4.

**BUG (BIOLOG Universal Growth agar) culture medium** contained 57.0 g BUG agar culture (BLG.70101, BIOLOG) and 1000 mL distilled water, pH 7.0–7.5.

**Positive Control Bacteria**

The positive control bacterium *Bacillus megaterium* As1,223, which serves as a P-solubilizing microbial fertilizer in common use (RodriGuez and Fraga, 1999; Chen et al., 2006), was supplied by the Microbe Collection Center of the Chinese Academy of Sciences (CAS).

### Isolation and Determination of the P-Solubilization Ability of Bacteria

The bacterial isolation protocol and determination of the P-solubilization ability were performed as previously described...
TABLE 1 | Physicochemical characteristics and nutrients in reclamation and experimental soil.

| Sites                | pH   | M.C (g/kg) | O.M (mg/kg) | Nitrate N (mg/kg) | Ammonium N (mg/kg) | Available P (mg/kg) | Available K (mg/kg) | Total N (mg/kg) | Total P (mg/kg) | Total K (mg/kg) |
|----------------------|------|------------|-------------|-------------------|-------------------|--------------------|---------------------|----------------|----------------|----------------|
| 4-year Reclamation   | 6.21 | 15.2       | 7.00        | 5.92              | 19.20             | 4.11               | 150.36              | 395.25         | 361.51         | 4625.84        |
| Experimental soil    | 4.0  | 22.3       | 26.54       | 8.12              | 29.35             | 6.39               | 355.90              | 676.22         | 480.36         | 5160.02        |

MC, moisture content; OM, organic matter.

(Chen et al., 2014). Briefly, for isolation of bacteria, each soil sample were homogenized in sterile distilled water and serially diluted. Aliquots of each dilution were spread on Nutrient agar medium and incubated at 30°C for 24–48 h. Colonies were selected on the basis of the development of a clear halo; the clones were further purified on Nutrient agar and TSB media. Once purified, each isolate was stored at −80°C in the same medium with 20% (v/v) glycerol.

For P-solubilizing ability determination, the isolates were screened by culturing at 28°C on the media supplemented either lecithin (Po culture medium) or Ca₃(PO₄)₂ (Pi culture medium). When the colonies appeared in 4–5 days, those causing a clear phosphate-solubilizing zone were selected out for further calculation. The size of phosphate-solubilizing zone was determined for each colony (Nair et al., 1995).

**Bacterial Colony and Mycelia Morphology Observation**

Candidate bacteria were inoculated on agar media and cultured for 48 h at 30°C. The bacterial colony shape and color were observed, and bacteria were Gram stained. The morphology and size of the bacterial samples were observed by scanning electron microscopy (SEM) (S-3400N, Hitachi) at the Beijing Agricultural Biotechnology Center.

**Taxonomical Assignment**

16S rRNA sequencing and phylogenetic analysis were performed for taxonomical assignment. The primers sequences are displaying as follows: 27 F: 5′-AGA GTT TGA TCC TGG CTC AG-3′, and 1492 R: 5′-TAC GGT TAC CTT GTT ACG ACT T-3′. Gene alignment was performed within the EzTaxon database, and a systematic phylogenetic analysis was performed using Mega 4.0 software. The phylogenetic tree was constructed by the neighbor-joining method with a bootstrap value of 1,000.

**Accession Numbers**

The nucleotide sequence data reported in this paper appear with the following accession numbers (GenBank: MN294691, MN305765, MN305766, MN305767, and MN305768).

**Gas Chromatography (GC)-Fatty Acid Methyl Ester (FAMES) Analysis**

Gas Chromatography-Fatty Acid Methyl Ester in each isolated PSB was determined using a Sherlock automatic bacterial identification system. All the identification procedures were completed by the Beijing Agricultural Biotechnology Research Center. TSB culture medium was applied for bacterial excitation. FAME profiles were then obtained by analyzing the samples on a GC (Agilent Technologies, United States) equipped with flame ionization detector (FID) and MIDI (R) Sherlock Microbial Identification System (MIDI Inc., Newark, DE, United States) software. FAMEs were identified according to their retention time, compared to a commercial standard mixture (MIS standard calibration, Part no. 1200-A) (Sasser, 1990).

**BIOLOG Identification**

The BIOLOG microbial ID system (Biolog Inc., Hayward, CA, United States) was used to further identify the physiological fingerprint of isolated bacteria strains, in accordance with the manufacturer’s manual. Briefly, the strains were cultured in BUG media for 24 h generations at 30°C and resuspended in inoculating solution. Then, 100 µL of culture was inoculated into each well of a Gen III MicroPlate and incubated at 33°C in the dark for 4–6 h or 16–24 h. The bacterial metabolites of PSBs were obtained using a spectrophotometer and then analyzed and compared with the BIOLOG database.

**Determination of the P-Solubilization Ability**

The S32 strain was cultured in Pi or Po media for 7 days with Ca₃(PO₄)₂, lecithin, CaHPO₄, powdered PR, FePO₄, or AlPO₄ as the P source. Non-inoculated medium was used as a blank control, while B. megaterium As1.223 was used as a positive control. The P-solubilization ability of the PSBs was tested three times independently. Each isolate was inoculated into a 200-mL conical flask containing 100 mL of Pi or Po liquid medium (as described above) and then shaken (160 rpm) at 30°C. The suspensions were sampled at the indicated days postincubation. To remove the insoluble culture media, the suspensions were kept stationary for more than 1 h. The acidity was assayed simply by reading on a pH meter, and the P availability was determined with the Molybdenum blue method (Watanabe and Olsen, 1965). Optimum pH (4–10), temperature (20, 25, 30, 35, and 40°C), carbon source (Glu, Suc, Sta, Fru, Lac, Man, and Gly), and sodium content (1–10%, w/v) for P-solubilization in liquid media were determined following the above method.

**Alkaline and Acidic Phosphatase Activity Detection**

Culture media supernatants from the lecithin-containing media were collected and detected using Alkaline and Acidic Phosphatase Kits (P0326 and P0321, Beyotime) in accordance with the manufacturer’s instructions. B. megaterium As1.223 and non-inoculated medium served as positive and negative controls, respectively. The phosphatase activity of 1 mg of phenol
production after a 15 min reaction with the matrix at 30°C in 100 mL of medium was defined as one King-Armstrong unit.

**Soil Experiment**

The S32 strain was cultured in 100 mL of TSB medium at 30°C for 36 h, and then cells were harvested by centrifugation at 5000 rpm at 4°C for 5 min and adjusted to $1 \times 10^8$ cells/mL in sterile deionized water by dilution and plating. The experimental and reclaimed soil were autoclaved, and the initial available P was about 6.41 mg/kg and 4.05 mg/kg. For the experimental group, 600 g of soil was soaked with the above S32 strain suspensions ($10^8$ cells/mL) for 30 min. Non-inoculated soil served as a negative control. The available P in the soil of each treatment group was determined at 10 and 30 days post administration, and the experiment was performed independently three times.

**Rice Seedling Growth Experiment**

Sterilized deionized water-washed 7-day-old Malaysia MR-219 rice seedlings were grown in experimental and reclaimed soil. For the experimental group, homogenous seedlings were soaked with S32 strain suspensions ($10^8$ cells/mL) for 30 min. Then, three seedlings were sown in each plastic pot. The plants were grown under natural growth conditions, maintaining a 20% absolute water content. Rice seedlings were harvested 21 days after sowing. Approximately $5 \times 10^8$/mL of live washed bacterial S32 cells were used as inoculum in each bacterial treatment.

The root morphology of the MR-219 rice was determined using a root scanner (Expression 1680, Epson). Total root length (cm), total surface area (cm²) and total volume (cm³) were quantified using a scanner (Expression 1680, Epson) equipped with a 2 cm deep plexiglass tank (20.30 cm) filled with H₂O (El Zemrany et al., 2007). The scanned data were processed by Win-Rhizo software (Regent Instruments Inc.).

**Organic Acid Evaluation**

Approximately 20 mL of the samples from each treatment were injected into HPLC with a UV detector set at 210 nm. A Rezex ROA-organic acid $\text{H}^+$ (8%) column was used, and the mobile phase was 0.005 N H₂SO₄ with a flow rate of 0.17 mL/min.

**Statistical Analysis**

All the data were processed with SPSS 13.0 statistical software and are presented as the mean ± standard deviation (SD). Student’s t-test analysis and one-way ANOVA was used to calculate the data variance, and $P < 0.05$ represents a significant difference.

**RESULTS**

**Screening and Isolation of the PSBs**

A total of 20 bacteria with a considerable transparent zone were screened and isolated by colony formation on Pi and Po plating media (Figure 1A). Among them, there were five bacteria with a diameter of the transparent zone (D) to colony (d) ratio larger than 1.50, as formed on Pi plating media, and a ratio of D/d on Po plating media larger than 1.90. Isolation S32 displayed the highest D/d ratio on Pi medium and Po medium, 4.24 and 5.19, respectively, (Figure 1B).

Five PSBs (H22, Y11, Y14, Y34, and S32) formed semicylindrical colonies that were emulsus, yellow, opaque, glossy, and orderly. SEM revealed that the thallus was rhabditiform, non-spore forming, and gram-negative with a size of 0.5–0.6 $\mu$m × 0.6–1.6 $\mu$m (Figure 2).

**Characterization of PSBs**

The DNA gene size of H22, Y11, Y14, Y34, and S32 was approximately 1.5 kb. The alignment of the 16S RNA of five PSBs was performed with the EzTaxon database, and a phylogenetic analysis was performed with highly homologous strains as reference bacteria. We found that Y14 was homologous to *Pantoea calida* 1400/07T (GQ367478) (Popp et al., 2010) (99.79%) (Figure 3A). H22, Y11, and Y34 were highly homologous to *Pseudomonas vancouverensis* ATCC 700688T (AJ011507) (Mohn et al., 1999), *Pseudomonas mandelii* CIP 105273T (AF058286) (Verhille et al., 1999), and *Pseudomonas frederiksbergensis* JAJ28T (AJ249382) (Andersen et al., 2000), respectively, (Figure 3B). S32 was highly homologous to *Pantoea* sp., of which *Pantoea rodasii* LMG 26273T (JF295053) (Brady et al., 2012) displayed the highest similarity (99.94%). S32 clustered with LMG 26273T and LMG 26275T in the phylogenetic tree (Figure 3C).
Stress Resistance Test of Five PSBs
Each PSB displayed high tolerance to a broad range of temperatures, pH values, and sodium concentrations, especially for S32. Its tolerance ranged from 4 to 37°C, pH values of 4–11, and sodium concentrations of 2–10% (Supplementary Table S1).

Fatty Acid Composition Analysis
Fatty acids were mainly distributed in the cell membrane as the basis of the lipids and lipopolysaccharides. The fatty acid composition data were analyzed according to the Microbial Identification System: the characteristic peak value >1% was
C_{12:0}, C_{16:0}, C_{17:0 cyclo}, and C_{18:1 \_7c} in the second and third peaks. The major characteristic peak contained C_{12:0} and C_{18:1 \_7c}, and the third peak occupied 27.6, 14.6, and 25.3% of that of the whole-cell component, which was consistent with that of LMG 26273\textsuperscript{T} and LMG 26275\textsuperscript{T} (Supplementary Table S2).

### BIOLOG Identification

To further characterize the S32 strain, which showed the highest P-solubilizing capacity among all the isolated PSBs, a BIOLOG test was performed to identify its species. The alignment results of strain S32 displayed good agreement with Pantoea cyri pedii at 22 h after culturing at 33°C (SIM value = 0.589). The BIOLOG reaction results ultimately confirmed that S32 was a Pantoea sp., which was consistent with our 16S rDNA sequence analysis (Supplementary Table S3).

### P-Solubilizing Efficiency of Five PSBs With Different P Sources

To screen out the bacterium with the highest P-solubilizing efficiency in Pi, Po, and powdered PR, the five selected bacteria were inoculated into 100 mL of liquid media supplemented with either Ca$_3$(PO$_4$)$_2$, lecithin, and powdered PR. For Ca$_3$(PO$_4$)$_2$, the P-solubilizing efficiency of S32 reached 24.19%, which was nearly quadruple that of the positive control B. megaterium As$_{1.223}$. The available P in the medium was 1256.67 mg/L, while the pH value was reduced by 3.51, which was double that of As$_{1.223}$ (Figure 4A). The P-solubilizing efficiency for lecithin was 2.47%, which was 1.5 times that of the positive control As$_{1.223}$. The available P content in the Po medium was 13.05 mg/L, and the pH value decreased by 3.89 (Figure 4B). However, with respect to PR, the five PSBs showed a degree of efficiency similar to that of As$_{1.223}$ (Figure 4C).

### P-Solubilization Capability of S32 Under Different Experimental Conditions

To determine the effect of inoculation time on the P-solubilization rate of S32, media containing Ca$_3$(PO$_4$)$_2$ was inoculated with S32 for different time points. The P-solubilization efficiency of S32 rapidly reached 23.91% at 24 h postinoculation (PI) and increased slowly to its maximum (24.61%) at day 5 PI. The content of available P in the medium reached 1204.67 mg/L at 24 h PI and then slowly peaked at 1332.33 mg/L at day 7 PI. The pH in the medium quickly decreased to 3.40 at 24 h PI, which was maintained for the following 6 days (Figure 5A).

The P-solubilizing efficiency of S32 on different P sources, Ca$_3$(PO$_4$)$_2$, CaHPO$_4$, lecithin, PR, AlPO$_4$, and FePO$_4$, was 24.09, 18.38, 2.47, 0.19, 0.51, and 2.62%, respectively. Each condition yielded a rate significantly higher than the B. megaterium As$_{1.223}$ positive control. The pH value decreased by 3.51, 2.68, 3.89, 3.58, 4.03, and 4.03 in these types of media, respectively. Thus, the efficiency of P-solubilization on Ca$_3$(PO$_4$)$_2$ was the greatest compared to the other P sources tested (Figure 5B). Notably, S32 showed significantly higher P-solubilizing efficiency than B. megaterium As$_{1.223}$ positive control.

By using glucose, sucrose, starch, fructose, lactose, mannitol, or glycerin as the carbon source, the P solubilization efficiency of S32 was 24.19, 0.75, 0.73, 8.29, 1.30, 7.40, or 5.25%, respectively, (the pH of the medium was reduced by 3.51, 0.23, 0.51, 1.82, 0.73, 2.23, or 1.56, respectively). This result demonstrated that glucose was the best carbon source for assisting the P-dissolving capacity of S32. Notably, we found that sucrose and starch did not differ from the negative control, suggesting that these two saccharides cannot be utilized as carbon sources by S32 (Figure 5C).

To assess the influence of pH on the P solubilizing efficiency of S32, the initial pH value of the Pi medium was set as 4.0, 5.0, 6.0, 7.0, 8.0, 9.0, or 10.0. After culturing for 24 h, the P-dissolving rate of bacteria S32 in these media was 13.56, 14.10, 22.32, 22.69, 22.51, 12.87, or 12.72%, indicating that the highest P solubilizing rate occurred at pH = 7 (Figure 5D).
Next, we determined the effect of temperature on the P-dissolving rate of S32. The P-dissolving rate for Ca$_3$(PO$_4$)$_2$ of bacteria S32 after it was cultured for 5 days at 20, 25, 30, 37, or 40°C was 21.13, 23.16, 24.19, 15.70, or 4.30% (the pH value of the medium was reduced by 3.15, 3.33, 3.51, 2.73, or 1.63), respectively. These data demonstrated that the best P-solubilizing efficiency was obtained at 30°C (Figure 5E).

The P-solubilizing ratio for Ca$_3$(PO$_4$)$_2$ by S32 in growth medium containing NaCl with concentrations of 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10% was 21.58, 20.35, 18.79, 18.57, 14.80, 13.85, 12.22, 10.79, 6.73, or 4.16% (the pH value in the medium was reduced by 2.90, 2.77, 2.58, 2.49, 2.70, 2.54, 2.37, 2.30, 1.87, or 1.13), respectively. These data suggested that the P-solubilizing capacity decreased with increasing sodium concentration (Figure 5F).

**Acid and Alkaline Phosphatase Activity of S32**

The activity of acid phosphatase (ACP) in S32 was 6.94 U/100 mL, which was 3.6-fold greater than the positive control As1.223
P fractions Before After Before After Before After
Av-P 4.05 12.89 4.02 6.90 4.00 4.05
Al-P 10.56 10.02 10.50 10.22 10.38 10.52
Fe-P 32.56 29.03 33.67 32.08 33.14 33.08
Oc-P 115.92 113.61 116.48 113.79 115.21 115.46
Ca-P 69.80 65.34 67.87 69.22 71.04 70.51
L-Po 9.00 8.36 8.83 9.36 9.03 8.76
ML-Po 19.54 17.02 18.60 16.31 19.00 19.05
MR-Po 14.22 11.02 14.45 13.34 14.30 14.24
HR-Po 51.60 46.44 50.62 49.88 51.22 51.06

Distributions of reclaimed soil P fractions before and after S32 inoculation. Av-P, available P; Al-P, Al-bound P; Fe-P, Fe-bound P; Oc-P, Occluded P; Ca-P, Ca-bound P; L-Po, labile organic P; ML-Po, moderately labile organic P; MR-Po, moderately resistant organic P; HR-Po, highly resistant organic P.

### DISCUSSION

In the present study, a total of five PSBs were screened from the reclaimed soils. These PSB isolates may possess the potential to be applied in improving soil recovery and crop production. A higher P-solubility capacity of S32 was observed in the Ca$_3$(PO$_4$)$_2$ medium compared to other bacteria and was therefore chosen for further investigation. Its P-solubilization rates for Ca$_3$(PO$_4$)$_2$ and lecithin reached 24.19 and 2.47%, which were nearly quadruple and double that of the positive control As$_1$.223, respectively. Furthermore, several insoluble P sources, such as CaHPO$_4$, lecithin, powdered PR, AlPO$_4$, and FePO$_4$ could also be dissolved by S32, with P-solubilizing rates of 18.38, 3.07, 0.16, 3.19, and 0.51%, respectively. Additionally, ALP and ACP could also be found in S32, with higher activities (6.94 U/100 mL and 1.223 U/100 mL) than the positive control B. megaterium As$_1$.223. Our study also suggested that S32 grew well and was adaptable due to its dependence on multiple carbon sources and tolerance to a broad range of temperatures, pH values, and sodium concentrations. It
also indicated that S32 achieved an optimal P-solubilizing rate in the following culture conditions: 5 days incubation, glucose as the carbon source, pH = 7, and temperature = 30°C.

Accumulating reports describe PSBs with the ability to dissolve both Pi and Po. For instance, Li et al. (2013) isolated four Pi-solubilizing bacteria from the plant *Anaphalis lutea*, and their P-dissolving rates ranged from 65.24 to 315.36 mg/L (D/d ratio of bacteria with the largest P-solubilizing cycle = 1.33). Wei et al. (2014) isolated two P-solubilizing fungi with rates of 1051.69 and 872.18 mg/L on Ca$_3$(PO$_4$)$_2$. Huang et al. (2010) isolated Pi-solubilizing bacteria with a Po-dissolving content of 537.6 mg/L and pH maximal decrease amplitude of 2.79. Lu et al. (2014) screened for Po-solubilizing bacteria with a D/d value of 4.3 and found that the available P in a medium with pH = 7 is 4.8 mg/L. However, the P-dissolving capability of both functions on reclaimed soil has rarely been reported. In the present study, the isolated bacteria S32 displayed higher P-dissolving capacity than the microbes in the above reports. The P-solubilizing cycles for Pi and Po were 4.42 and 5.19, respectively, while the available P in the Pi and Po media were 1256.67 mg/L and 13.05 mg/L, respectively. Remarkably, we tested the P-solubilizing efficiency of S32 in Xiaoai reclaimed soil, and the data showed that S32 also displayed a considerable P-solubilizing efficiency.

**TABLE 5** | Effect of S32 on the root growth of plants in ES or RS.

| Soil        | Root length (cm) | Root surface area (cm$^2$) | Root volume (cm$^3$) |
|-------------|------------------|---------------------------|---------------------|
| S32         | 57.21 ± 3.56*    | 56.63 ± 9.68**            | 5.99 ± 0.25         |
| PC          | 34.52 ± 6.25     | 28.45 ± 5.59              | 4.50 ± 0.24         |
| NC          | 25.81 ± 5.65     | 25.14 ± 8.50              | 3.96 ± 0.85         |

*ES, experimental soil; RS, reclaimed soil; PC, positive control; B. megaterium As1.223; NC, negative control. *P < 0.05; **P < 0.01 compared to NC.*

Phosphate solubilizing bacteria mainly belong to *Bacillus* (Babu et al., 2017; Thomas et al., 2018), *Pseudomonas* (Otieno et al., 2015; Paul and Sinha, 2017), *Arthrobacter* (Jiang et al., 2019), *Agrobacterium* (Rodriguez and Fraga, 1999), *Micrococcus* (Dastager et al., 2010), *Enterobacter* (Jiang et al., 2019), *Vibrio* (Yu et al., 2012), *Serratia* (Behera et al., 2017), *Rhizobium* (Korir et al., 2017), *Aeromonas* (Jha et al., 2012), and *Burkholderia* (Jha et al., 2012). Several fungi display the same function, including *Sclerotium* (Thampi and Bhai, 2017), *Penicillium* (Li et al., 2016), *Aspergillus* (Yin et al., 2017), and *Trichoderma* (Lei and Zhang, 2015). Huang et al. (2010) found that *Pseudomonas* and *Pantoea* are optimal species for their stable P-solubilizing effects. Son et al. (2006) isolated a P-dissolving *Pantoea* from the soybean rhizosphere, and soluble P in its culturing medium reached 900 mg/L. Zhang et al. (2013) isolated another *Pantoea* from the *Caragana microphylla* rhizosphere, and rapidly available P in its Pi culture media reached 4.45 mg/L. Kaur and Reddy (2013) isolated efficient P mineralizing bacteria and tested their efficacy in plant mineral uptake and soil fertility of an organic field. Amongst 12 PSB isolated from an organic field, two isolates were selected for field inoculation based on their RP solubilizing ability, exudation of organic acids, phosphatase and phytase activity and production of indole acetic acid and siderophores. These isolates were identified as *Pantoea* cypripedii and *Pseudomonas* plecoglossicida. These isolates significantly increased yield and total P uptake in maize. Soil analysis showed that available P, organic carbon and soil enzyme activities were significantly increased (Kaur and Reddy, 2013). Castagno et al. (2011) isolate 50 PSB strains from a constrained environment such as the Salado River Basin in Argentina. Subsequently, they were found to be related to *Pantoea, Erwinia, Pseudomonas, Rhizobium* and *Enterobacter genera*, via 16S rRNA gene sequence analysis. The most efficient isolate, was identified as *Pantoea* eucalypti, a novel species in terms of plant growth-promoting rhizobacteria (Castagno et al., 2011). Park et al. (2011) isolated 18 different PSB strains from P amended and Lead (Pb) contaminated soils were screened for their efficiency in P solubilization. One PSB was chosen for Pb immobilization and was identified as *Pantoea* sp. The PSB significantly increased P solubilization by 49.9% in the case of *Pantoea* sp. for 800 mg/kg of RP addition, respectively, thereby enhancing the immobilization of Pb by 8.25–13.7% (Park et al., 2011). These reports strongly suggested *Pantoea* sp. a potential candidate for highly efficient PSB. In the present study, the soluble P content in S32 medium reached up to 1256.7 mg/L.

**TABLE 6** | Mean leaf and root P concentrations of plants inoculated with S32 and controls in ES or RS.

| Treatment | Leaf P conc. in ES (µg P/mg) | Root P conc. in ES (µg P/mg) | Leaf P conc. in RS (µg P/mg) | Root P conc. in RS (µg P/mg) |
|-----------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| S32       | 200.23 ± 29.13*               | 45.12 ± 5.67**                | 123.41 ± 15.51*               | 36.36 ± 3.02*               |
| PC        | 143.21 ± 10.46                | 31.95 ± 3.69                  | 80.24 ± 12.32                 | 28.25 ± 2.56                |
| NC        | 103.42 ± 11.39                | 24.19 ± 7.90                  | 69.15 ± 9.54                  | 22.96 ± 1.62                |

*ES, experimental soil; RS, reclaimed soil; PC, positive control; B. megaterium As1.223; NC, negative control. *P < 0.05, compared to NC.*

**TABLE 7** | Effect of S32 on the organic acid release in ES.

| Treatment | Oxalic acid (mM) | Citric acid (mM) | Malic acid (mM) |
|-----------|------------------|------------------|-----------------|
| S32       | 93.25 ± 4.02*    | 52.66 ± 5.88     | 83.14 ± 14.10** |
| PC        | 78.14 ± 6.84     | 51.20 ± 5.08     | 5.26 ± 3.68     |
| NC        | 69.00 ± 4.03     | 25.35 ± 2.31     | 9.25 ± 2.75     |

*ES, experimental soil; RS, reclaimed soil; PC, positive control; B. megaterium As1.223; NC, negative control. *P < 0.05; ***P < 0.001 compared to NC.*
demonstrating that S32 possesses a high P-solubilization capacity, even compared to other strains of Pantoea.

Over 80% of the P in cropland soil is Pi. Furthermore, Pi components in various types of soil display significant differences, e.g., calcium and magnesium phosphate are dominant in some types of soil, while ferric and aluminum phosphate are more abundant in others (Yu et al., 2012). In the present study, the aluminum phosphate-dissolving rate of S32 was 145.95 mg/L, which was lower than that of the PSB reported by Wang et al. (2013). However, S32 was able to dissolve multiple insoluble P sources other than aluminum phosphate. Different levels of soluble phosphates in various forms of P sources affect the solubilization of insoluble phosphates. First, the process of solubilization by the bacteria was documented to be regulated by the external phosphate levels, as also observed for phosphatases (Nahas, 2007). ACP production was reduced while concentration of soluble phosphate was increased (Wang and Lambers, 2019). Second, soil fertilization with P is important to both plant and microorganism growth. Increasing active exudation from roots provides substantial amount of organic acids that enhance solubilization. After soluble phosphate was exhausted, the solubilization process was triggered, with a consequent increase in soluble phosphate in the soil (Nahas, 2007). In the present study, the P-solubilizing efficiency of S32 on different P sources, Ca₃(PO₄)₂, CaHPO₄, lecithin, PR, AIPO₄, and FePO₄, was 24.09, 18.38, 2.47, 0.19, 0.51, and 2.62%, respectively.

As heterotrophic bacteria, phosphate solubilizers required carbon source and energy for both the synthesis of new cell material and the oxidation of carbon compounds. Rhizosphere soils present water-soluble C compounds mainly as carbohydrates and organic acids and a small portion and amino acids. It was well known that increasing number of microorganisms was associated with the plant rhizosphere due to its carbon concentration (Nahas, 2007). In the present study, by using glucose, sucrose, starch, fructose, lactose, mannitol, or glycerin as the carbon source, the P solubilization efficiency of S32 was 24.19, 0.75, 0.73, 8.29, 1.30, 7.40, or 5.25%, respectively.

S32 inoculation increased plant height and biomass. In addition to P solubilization activity, PSB was reported to secrete phytohormones that might have an influence on root growth. The extensive root system increased nutrient uptake from the surroundings, which increased plant biomass (Wang and Lambers, 2019). Soluble organic acids could serve as a source of carbon for microorganisms and subsequently affect the rhizosphere microbial environment, as well as plant growth. The plant root development in rice was affected by the application of PSB. Therefore, we also tested the effect of S32 on rice root growth and organic acid production in experimental and reclaimed soil.

A plant with S32 inoculation was found to possess fast root growth and release high amounts of organic acids. These results are consistent with the findings of Srivastava et al. (2007) who reported that the addition of PSBs resulted in the release of P and positively affected plant growth. Moreover, Hoffland et al. (1989) found that increased organic acids in the rhizosphere microbial environment were highly efficient at increasing the P-solubility of PSBs. The root development and plant biomass were highly correlated with the higher availability of P; in addition, PSB application may also have some other beneficial effects, such as phytohormone production.

**CONCLUSION**

In conclusion, the bacterium Pantoea sp. S32, possessing high dissolving capacity for both Pi and Po, was isolated from alfalfa rhizosphere soil in the reclamation area. S32 showed remarkable P-solubilization rates for different P sources and promoted plant growth, suggesting that the present study provides a potential approach for accelerating minefield recovery and increasing crop output in reclaimed soil.

**AUTHOR CONTRIBUTIONS**

QC performed all the experiments, and collected and analyzed the data. SL conceived the idea, analyzed the data, and wrote the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2019.02171/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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