



Nutrient Uptake in Cowpea Improved By Plant Growth Promoting Bacteria

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ABSTRACT

Abstract: Plant growth promoting bacteria including *Agrobacterium* spp, *Azotobacter* spp, *Pseudomonas* sp. and *Paenibacillus* s., which are known to promote the growth of plants were used to inoculate cowpea alongside *Rhizobia* to determine their effects on the uptake of nutrients such as Nitrogen, Phosphorus, Calcium, Sodium, Magnesium and Potassium which all play specific roles in the development of plants and are important for their growth. The experimental cowpea plants were allowed to grow for eight weeks under screen house conditions during which Nitrogen-free nutrient was applied to them weekly. After harvesting, the quantity of these nutrients in the dried shoot was determined. It was observed that the consortium of *Agrobacterium* spp, *Azotobacter* sp. and *Paenibacillus* sp improved the uptake of nitrogen and phosphorus, a consortium of *Pseudomonas* sp. and *Paenibacillus* sp. improved the uptake of nitrogen, magnesium and calcium while the consortium of *Azotobacter* spp improved the uptake of calcium. There was no specific pattern in the uptake of sodium and potassium.

Keywords: Nutrient uptake, Plant growth promoting bacteria, Cowpea

iSTEAMS Cross-Border Conference Proceedings Paper Citation Format

Ajayi, O. O., Dianda, M., Fagade O. E.* & Nwadike, B. (2018) Nutrient Uptake in Cowpea Improved By Plant Growth Promoting Bacteria. Proceedings of the 13th iSTEAMS Multidisciplinary Conference, University of Ghana, Legon, Accra, Ghana. Vol. 2, Pp 255-262.

1. INTRODUCTION

Nitrogen fixing bacteria especially members of the family Rhizobiaceae have been known to be able to fix nitrogen when they are found to be in friendly symbiotic relationships with leguminous plants. This phenomenon is described as nitrogen fixation and is known as nature's nitrogen factory (13). Although much work has been done on legume – rhizobium symbiosis, it has been found that other microorganisms are also important and useful roles in nitrogen fixation. These other endophytic bacteria which can be found in legume nodules are able to penetrate the nodules alongside rhizobia but they have been ignored for a long time (1,12) there is therefore little information on the roles they play in enhancing plant growth especially with respect to increased absorption of macronutrients such as phosphorus and their importance as plant growth promoting rhizobacteria (PGPR). Nitrogen (N₂) is an essential nutrient required for plant growth as it is a prominent component of amino acids, nucleic acids and chlorophyll. Deficiency of nitrogen results in chlorosis which is characterized by yellowing of leaves, stunted growth, slow growth, etc. Despite its abundance in nature, N₂ is unavailable to plants because of its inert nature.

Phosphorus is a fundamental component of the substances that are the building blocks of genes and chromosomes (14,19). It is an essential part of the process of carrying the genetic code from one generation to the next, giving the blueprint for all characteristic of plant growth and development (16,17). Sufficient phosphorus is also required to enhance different plant organs growth, promote nodulation and early maturity in legumes (6). Potassium is required for every major step of protein synthesis. The process of the "reading" the genetic code in cells of plants to produce proteins and enzymes that regulate all growth processes would be impossible without adequate K. When plants are deficient in K, there is no synthesis of proteins in spite of an abundance of available nitrogen (N). Instead, there is accumulation of protein "raw materials" (precursors) such as amino acids, amides and nitrate. The enzyme nitrate reductase catalyzes the formation of proteins, and K is likely responsible for its activation and synthesis (7).

Calcium (Ca) is taken up by plants as the calcium ion (Ca⁺⁺). A structural nutrient, it is an essential part in cell walls and membranes and is required for the formation of new cells. For this reason, early season availability of supplemental Calcium has a distinct effect on fruit set. Once deposited in plant tissues, Calcium is not remobilized. Therefore, young tissue is affected first under conditions of deficiency. Since Calcium is not mobile, the requirements of a crop for Ca⁺⁺ after early fruit set are commonly supplied in the form of nutritional sprays (14). Plant uptake of magnesium is in the form of the magnesium ion (Mg⁺⁺).



The chlorophyll molecule, which is essential for photosynthesis, contains magnesium (19). Sodium is not an essential requirement for plants but is used in small quantities by plant where it affects photosynthesis by altering the balance of water, aids metabolism and synthesis of chlorophyll. Its deficiency does not show any symptom in plants (13,15).

Materials and methods

2.1 Study area and sample collection

Samples were collected from three sites in Nassarawa State which are Shama Local government (N 08° 37' 47.7" E 007° 46' 48.4" Elevation 244 m), Ogba/ Ube Egon local government (N 08° 51' 55.4" E 080° 25' 34.5" Elevation 399 m), Mandara Kokona local government (N 08° 50' 29.8" E 008° 12' 37.1" Elevation 364 m). Nassarawa State was selected because at the outset of this research the nodule samples were readily available and it is also an area where large cultivation of cowpea is a common practice. A hand trowel was used to mark a circle with a radius of approximately 15 cm around the base of plant, and section was cut out to a depth of about 20 cm. A spade was then used to slowly lift out the clump. The soil particles were carefully removed from the root material mechanically. Detaching of the secondary roots from the plant was avoided as nodules may be found on the lateral roots as well as the tap root. A sieve of 0.5 mm size and mesh was then placed under each root sample to catch nodules that may become detached from the root. The root was then carefully washed under a gentle stream of water from a tap in a bowl in a sink (11). The samples were then wrapped in Aluminum foil paper and transported by road to Ibadan.

2.2 Isolation of Rhizobia from Root Nodules

Microorganisms were isolated from nodules on Congo red agar (20) using spread plate method. Five undamaged nodule samples were selected at random from each site. They were placed in sterile water for about 15 to 20 mins to rehydrate them after which they were surface sterilized using 3 % sodium hypochlorite for 3 minutes. They were then rinsed in sterile water; then further sterilized with 95 % ethanol and rinsed with six changes of sterile water. The nodules were transferred into sterilized petri-dishes and crushed with flamed glass rod. A few drops of sterile water were added to the crushed sample. A loop full of crushed nodule was streaked on Congo red agar and then incubated at 28°C for 5 - 7 days (18, 20). Selected isolates present were picked, rhizobial isolates were selected based on their cultural appearances i.e. their ability to absorb Congo red dye thus appearing white on the Congo red media (18, 20). Isolates were labelled, purified and stored on yeast mannitol agar slants and nutrient agar slants. The major media used for isolating and culturing of rhizobia were Congo red, Yeast Mannitol Broth, and Nutrient agar.

2.3 Pot Experiment

Pure cultures of the rhizobia isolates were obtained and introduced into 100 ml Erlenmeyer flasks containing 50 ml of yeast-mannitol broth in duplicates. The pure cultures of the Non rhizobial microorganism which were to be co-inoculated with the rhizobial isolates were also obtained and introduced into one of the duplicate rhizobial broths. For Rhizobial isolates (R1, R2, R3) obtained from Shama, the following plant growth promoting rhizobacteria (PGPR) were added: *Agrobacterium spp.*, *Azotobacter sp.*, and *Paenibacillus wynnii* (consortium A) while the Rhizobial isolates obtained from Mandara (R4,R5) were co-inoculated with *Paenibacillus wynnii* and *Pseudomonas aeruginosa* (consortium B) and the rhizobial isolates from Ogba (R1, R2, R3,R4,R5,R6, R7, R8, R9) were co-inoculated with two *Azotobacter spp.* (consortium C). The inoculated broth was incubated at room temperature (28°C) on a Rotary shaker for 7 days and then used to inoculate plants at 1 week of growth (20). Sea sand was collected from Lagos beach and washed with water to remove debris and pH till water is clean and clear. The crushed gravel and medium sized gravel were also washed till the water was clean. The sea sand, crushed gravel and peat were mixed in a ratio 6:6:1 and mixed until it was evenly distributed. The mixture was then sterilized at 121°C and 1.05 kg cm⁻² for 15 minutes. The medium sized gravel was also sterilized (20). 500 ml pots were washed and sterilized using 5 % of JIK (3 % w/v sodium hypochlorite) after which it was thoroughly rinsed twice with sterile water to ensure complete removal of all traces of sterilizing agent. Seeds were planted in 500 ml pots and allowed to germinate. One week after planting (WAP), the cowpea plants were thinned to remain one viable plant per pot after which pots were labelled. 1 ml of the inoculums which were already prepared as described above, was introduced into the one week old cowpea plants using sterile pipette. A total of 18 treatments were used in addition to Nitrogen controls (N+ treatment i.e. to which nitrogen was applied).

2.4 Application of nutrient to Plants

Cowpea plants were allowed to grow for 8 weeks during which they were given 20 ml of nutrient solution consisting of both micro and macro nutrient. To prepare nutrient solution given to plant, the stock solutions were mixed using 100 ml of macro- stock solution and 10 ml micro- stock solution made up to 10 liters using distilled water. The nutrient solution was sterilized at 121°C and 1.05 kg cm⁻² for 15 minutes and was aseptically given to the plants weekly. The solution for the N+ treatment (control containing nitrogen) was prepared using 5% of N in KNO₃ this was sterilized at 121°C and 1.05 kg cm⁻² for 15 mins after which 50 ml of the solution was added to the plant weekly.



2.5 Harvesting

The plants were harvested 8 weeks after planting (WAP) by cutting at the base with a secateur. The shoots and roots were collected and placed in labeled paper bags and placed in an oven where they were dried at 68°C for 72 hours until constant weight was obtained.

2.7 Determination of nutrients in plant shoots

The total Nitrogen in plant shoots was determined using the micro Kjeldahl method (8) phosphorus was determined by the molybdenum blue method as described by (10). The concentrations of Ca, K, Na and Mg in shoot were determined following the method described by (5). Unnodulated plants of the cowpea N⁺ treatment was used as a reference plant

3. RESULTS

3.1 Nitrogen

Plant samples were processed and analyzed for total-N. The total yield from the percent total-N in the plant material and the dry matter yield were calculated, and the difference in total-N yield between the cowpea and the reference crop obtained, to give the estimate of N₂ fixation. Total Nitrogen for at R1, R2, R3, R4, R5 on addition of consortium A and B respectively were higher than those that had only the rhizobial inoculation, while R6, R7, R8, R9 to which *Azotobacter* spp. had lower Nitrogen value than their counterparts to which only rhizobia strains were added. All treatments gave a higher N value than that of the N⁺ treatment with a percentage increase ranging between 7.8 – 50.9 % See Fig 1.

3.2 Phosphorous

The rhizobia strains to which consortium A was added (R1, R2, R3) had a higher P value than their counterparts with only rhizobia while R5 and R6 to which consortium B and consortium C were added respectively also had higher P values with only rhizobial inoculants although others in their group showed a negative response. About 60 % of the treatments had a higher P value than N⁺ treatment (See Fig 2).

3.3 Calcium

All treatments had a higher Ca value compared to the N⁺ treatment. Consortium i.e R6, R7, R8, R9 to which *Azotobacter* spp was added showed consistency in improving the uptake of calcium when compared with their counterparts to which only rhizobia was added. While R2 and R5 also had higher values of calcium uptake the other in their group did not

3.4 Potassium

All treatments had a lower K value than N⁺ because KNO₃ was used as a source of nitrogen for the N⁺ treatment thus a higher amount of Potassium was available to the N⁺ plants. Fig 3

3.5 Sodium

All treatments had a lower Na value than the N⁺ (See fig 6) including those to which the three groups of PGRBs were added. There was also no consistent pattern in the absorption of sodium suggesting that these group of PGRBs including *Agrobacterium* spp, *Paenibacillus wynnii*, *Pseudomonas aeruginosa* and *Azotobacter* spp did not have any effect on the uptake of sodium but that other factors or PGRBs may be responsible for its uptake. See fig 6

3.6 Magnesium

Consortium B i.e treatments R4, R5 to which they were added were able to improve the uptake of magnesium while the consortium of A and C showed no consistency in increasing the uptake of magnesium. 83.3% of the treatments had a higher Mg value than the N⁺ treatment, While R2, R7 and R9 also showed increased the Mg content on addition of PGRB compared to the others in their group obtained are shown in fig 5.

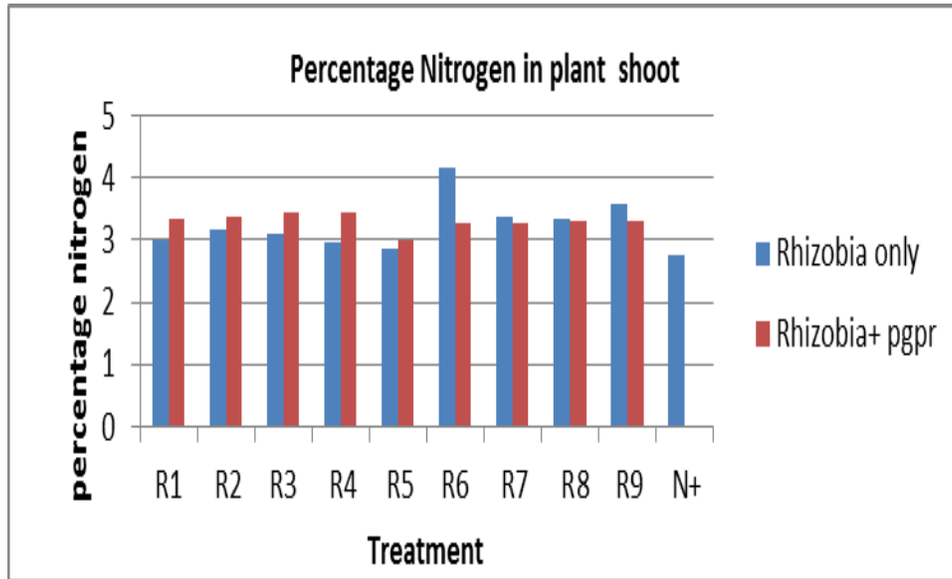


Fig1: Percentage Nitrogen in the dried shoot of plant

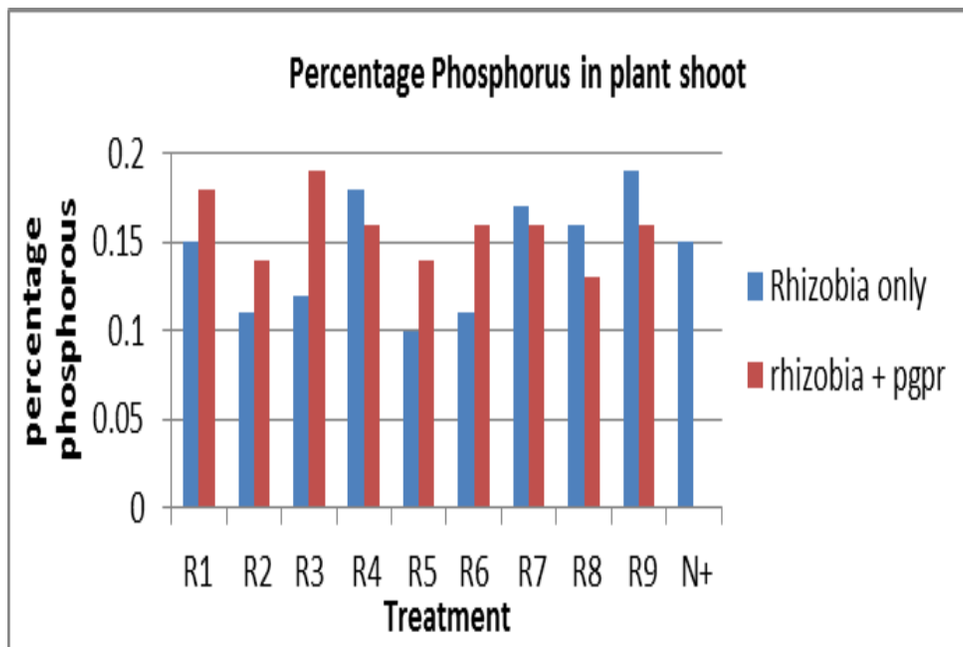


Fig 2: Percentage Phosphorus in the dried shoot of plant.

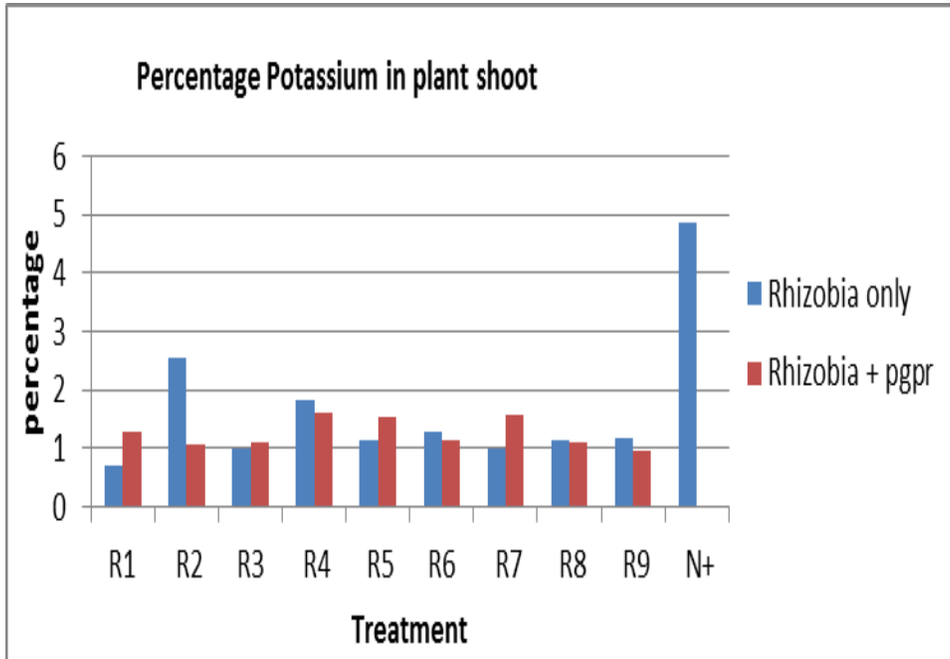


Fig 3: Percentage Potassium in the dried shoot of plant.

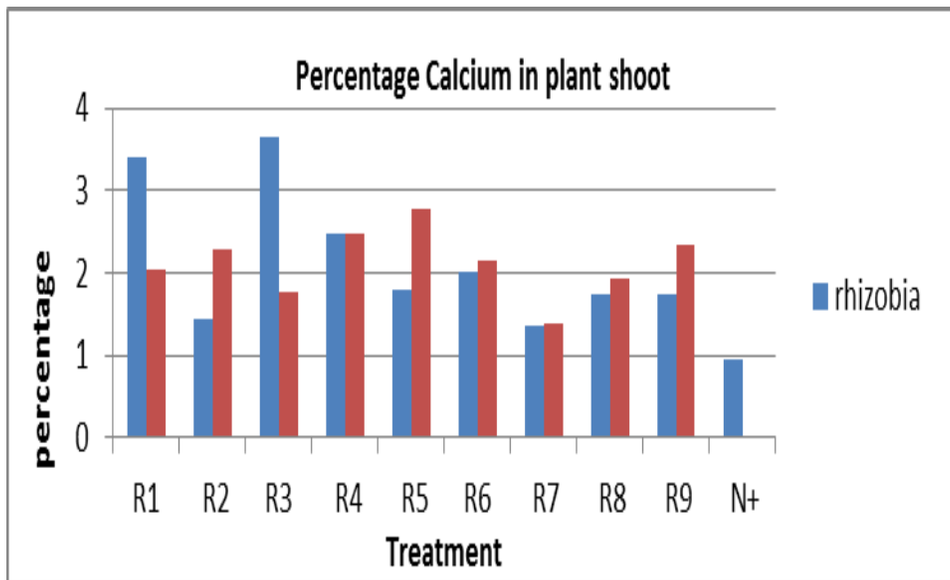


Fig 4: Percentage Calcium in the dried shoot of plant

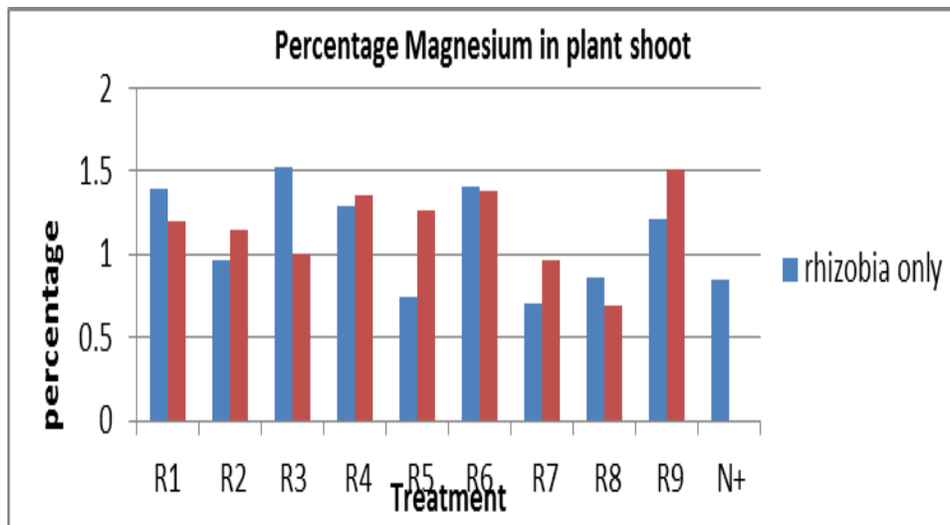


Fig 5: Percentage Magnesium in the dried shoot of plant

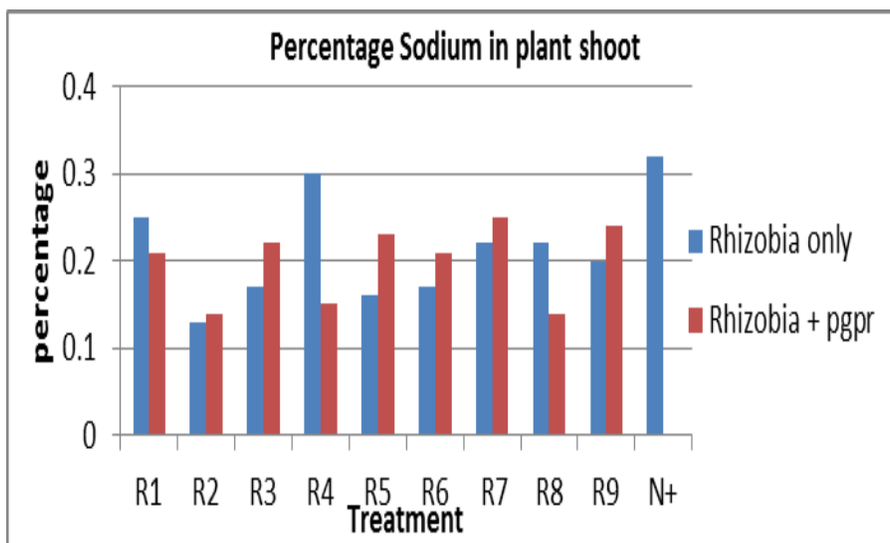


Fig 6: Percentage Sodium in the dried shoot of plant

4. DISCUSSION

This research was carried out to determine if Plant growth promoting bacteria isolated from the nodules of cowpea can increase the absorption of some macro nutrients including Nitrogen, Phosphorous, Magnesium Calcium, Potassium and Sodium. Total Nitrogen for R1, R2, R3, R4, R5 on addition of PGPBs was higher than those that had only the rhizobial inoculation, and all treatments had a higher N value than that of the N⁺ treatment this is similar to the work of Eutropia *et al.*, (2014) (where there was significant uptake of nutrient was observed in experiments using soybean), the work of Ahmad *et al.*, (2008) and Young *et al.*, (2001). All rhizobial had a higher P value on addition of PGPRs except R9 and R8. About 60% of the treatments had a higher P value than N⁺ this was similar to the result obtained by Eutropia *et al.*, (2014) where significant uptake of P was observed.



All treatments had a lower k value than N^+ because KNO_3 was used for the N^+ treatment thus a higher amount of Potassium was available to the plants absorption of Na showed no particular pattern and all treatments had a lower Na value than the N^+ treatment This suggests that these group of PGRBs including *Agrobacterium* spp, *Paenibacillus wynnii* *Pseudomonas aeruginosa* and *Azotobacter* spp. did not have any effect on the uptake of sodium but that other factors or PGRBs may be responsible for its uptake a similar report was given by Mehrpouyah *et al.*, (2012) and Eutropia *et al.*, (2014). All treatments had a higher Ca value compared to the N^+ treatment. Consortium i.e R6,R7,R8,R9 to which *Azotobacter* spp. were consistently able to improve the uptake of calcium when compared with their counterparts to which only rhizobia was added as was also reported in work of Makoi *et al.*, (2013) and Yahya-Abadi; (2008).For magnesium, all treatments had a higher percentage of magnesium than the N^+ treatment except R5, R7, R8 co-inoculated with *azotobacter* spp while consortium B increased the uptake of magnesium consistently. Rhizobia strain 5 showed ability to work with all the PGRBs used by increasing the uptake of all the nutrients after it was co-inoculated with them.

Conclusion: Consortium A (*Agrobacterium* spp, *Azotobacter* sp., and *Paenibacillus wynnii*) when added along-side rhizobia strains were able to increase the uptake of nitrogen and phosphorous, while Consortium B (*Paenibacillus wynnii* and *Pseudomonas aeruginosa*) increased the uptake of nitrogen, magnesium and calcium and Consortium C (*Azotobacter* spp.) increased the uptake of calcium.

Acknowledgement: This work was supported by the International Institute of Tropical Agriculture Ibadan, Oyo State Nigeria.

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