Influence of microbial priming on germination and seedling growth traits of compact cotton CO17

ABSTRACT

Cotton, known as "the King of fibers", is the predominant fiber in the Indian textile industry. Plant growth-promoting rhizobacteria (PGPR) represent a potential sustainable alternative for the enhancement and protection of crops. The germination and seedling growth of cotton can be optimized by inoculating with PGPR. An experiment was conducted to evaluate the effect of different PGPR strains on seed germination and seedling establishment characters on cotton. The highest germination percentage, maximum vigour index and leaf area was obtained with the *PPFM* TNAU1 strain inoculation. The maximum shoot and root length were observed with seeds treated with *Azospirillum* strain sp7 with an increase of 24.4 and 42.8,% over the control. Underground fresh and dry matter was recorded higher in the seedlings treated with *Azospirillum* sp7 strain compared to control, while the *PPFM* TNAU1 strain treatment increased the aerial fresh and dry matter content because of its larger leaf area. Seeds inoculated with individual strain of *Azospirillum* sp7 and *PPFM* TNAU1 outperformed the combined inoculation of PGPR strains. The increase in germination traits and seedling characters by PGPR strains indicates the positive influence on improving cotton seedling establishment traits associated to for higherbetter yield.

Keywords: Cotton, plant growth promoting rhizobacteria, microbial priming, germination and seedling emergence traits.

1. INTRODUCTION

Cotton is one of the predominant commercial and a crops with ef-global significance, playing an important—momentous role in foreign exchange and industrial economy. Current production of cotton fiber is not sufficient to meet the gradually-increasing demand of world population which arises due to many limiting factors, *i.e.*, frequent droughts, soil degradation—of soils, salinity, and alkalinity. To date, many farming practices were employed to increase cotton production, as a consequence, toxic impacts were implied on water and soil resources. In addition—on top of that, a major part of the chemical fertilizers applied to the crops remain in the soil as insoluble inorganic compounds which and promotes soil toxicity—to the soil. Lately, efforts have been focused on minimizing the useage of chemical fertilizers in order to optimize cost of—production cost—and protect the environment against pollution without compromising the seed cotton yield. The use of biological stimulators in Indian agriculture has many economic and ecological advantages. The search for alternative solutions has stimulated prompted researchers to take a second look at the range of microorganisms which provide benefits to agricultural production by stimulating plant growth and producing—a higher yield [1,2].

Plant growth promoting rhizobacteria (PGPR) are the bacteria living in the rhizosphere region which interacts with plant metabolism and improving advances—their growth directly or indirectly. Newadays, PGPR inoculants are considered as a part of integrated nutrient management system to improve plant growth and development. The use of beneficial bacteria such as Azotobacter, Azospirillum, Acetobacter, Pseudomonas, Methylotrophs, Bacillus, pPhosphobacteria etc. colonizes plant root and promotes growth through by nitrogen fixation, phosphorus and potassium solubilization. Seed priming by the microbial inoculants favours increased germination by the activating on of germination related enzymes, increasing metabolism that helps in the rapid growth of radicle and plumules. These PGPRs helps in improve better—seed germination and colonizes plant roots to improvinge seedling vigour, and modifies root morphology to facilitate better acquisition of nutrients increasing which ultimately enhances the crop yield and quality of crop.

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Higher production and productivity of the crop is achieved through the use of good quality seeds and

2. MATERIAL AND METHODS

2.1. Experimental site and soil characteristics

The field experiment was conducted at Eastern block farm, Tamil Nadu Agricultural University, Coimbatore, located in the Western Agro-climatic zone of Tamil Nadu (11° 02' N latitude, 76° 93' E longitude, and at an altitude of 428.5 masletres above mean sea level). The experiment was planted laid out following in thea randomized block design with four replications. Experimental soil was sandy clay loam with pH 8.49 and organic matter content 0.47%. Newly released compact cotton seeds (CO 17 cultivarvariety) seeds were obtained collected from the Cotton Department of Cotton, Tamil Nadu Agricultural University, Coimbatore. Planting Sewing was done in rows by dibbing at row to row 90 cm apart and plant to plant 15 cm spacing by placing the seeds at each hill. Irrigation was given as per requiredment of crop. Standard plant protection schedule was followed to protect the crop from diseases and pests as per recommended package of practices of Tamil Nadu Agricultural University, Coimbatore.

2.2. Seed treatment details

Four PGPR strains included i.e., Azospirillum sp7, Phosphobacteria PS1, Potash releasing bacteria KRB9 and Pink pigmented facultative methylotrophs TNAU 1 were obtained collected from the Department of Agricultural Microbiology, Tamil Nadu Agricultural University, Coimbatore. The treatments include T1 - Control, T2 - Azospirillum sp7 (50 ml/acre of seeds), T3 - Pink Pigmented Facultative Methylotrophs (PPFM) TNAU 1 (50 ml/acre of seeds), T4 - Azospirillum sp7 + Phosphobacteria PS1 + Potash releasing bacteria KRB9 (50 ml/acre of seeds), T5 - PPFM TNAU1+ Phosphobacteria PS1 + Potash releasing bacteria KRB9 (50 ml/acre of seeds), T6 - PPFM TNAU1+ Phosphobacteria PS1 + Azospirillum sp7 + Potash releasing bacteria KRB9 (50 ml/acre of seeds). Azospirillum, PPFM, Phosphobacteria and Potash releasing bacteria are the commercial formulations of biofertilizers which contains Azospirillum brasilense sp7 strain, PPFM TNAU1 strain, Bacillus megaterium PS1 strain and Bacillus mucilaginosus KRB9 strain, respectively. Cotton seeds were bioprimed using 2% CMC (Carboxymethyl Cellulose) solution and then—shade dried before plantingprior to sowing.

2.3. Data collection

The efficiency of different PGPR strains as bio_-inoculants as an individual and interactive effect on seed germination efficiency, seedling emergence and growth-traits in cotton were recorded.

2.3.1. Germination percentage - GP (%)

Final germination percentage was calculated by dividing the number of seeds germinated by the total number of seeds and expressed in % [3].

2.3.2. Mean germination time - MGT (day)

Mean germination time (MGT) was given by [4].

 $MGT = \sum \left(\frac{ni*ti}{ni}\right)$

where ni is the number of germinated seeds on germination days, ti is the number of days during the germination period (between 0 and 10 days)

2.3.3. Germination rate index - GRI (Germination % day 1)

The germination rate index was computed by the formula given by [5]

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$$GRI = \frac{G1}{1} + \frac{G2}{2} + \frac{GX}{X}$$

where G1=Germination percentage at the first day after sowing, G2= Germination percentage at the second day after sowing.

2.3.4. Coefficient of velocity of germination - CVG

The coefficient of velocity of germination was calculated by the formula given by [6]

$$CVG = \frac{\sum ni}{\sum ni*ti} * 100$$

where ni is the number of germinated seeds on day ti; ti = the number of days during the germination period (between 0 and 10 days)

2.3.5. Germination index - GI

The germination index (GI) was calculated based on the formula of [7]

$$GI = (10 * n1) + (9 * n2) + \dots + (1 * n10)$$

where n1 = number of seedlings emerging on first day after planting; n2 = number of seedlings emerging on second day; n10 = number of seedlings emerging on tenth day.

2.3.7. Seedling vigour index - SVI

The vigour index value was <u>obtained computed</u> by multiplying germination of seeds in percentage and total seedling length in centimeter and expressed in whole number as described by [8]

2.3.8. Root length and shoot length (cm)

Root and shoot length were measured at 10 days old seedlings and expressed in centimeters (cm)

2.3.9. Leaf Area (cm²)

Leaf samples collected from each replication were cleaned well—and inserted into_a leaf area meter (LICOR, Model LI 3000) and leaf area measured was expressed as cm² per plant.

2.3.10. Aerial and underground weight (g seedling-1)

Fresh aerial and underground weight of 10 days old seedlings were taken and expressed in g seedlings. The sSeedlings used for growth measurement were placed in a paper cover and dried in shade for 24 hrs and then they were placed kept in a hot air oven at 65 °C for 48 hrs. The dried seedlings were weighed to estimate the aerial and underground dry matter production and the mean values were expressed in mg seedlings. 1.

2.4. Statistical analysis

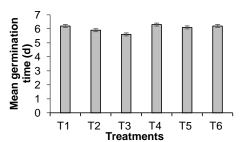
Data were analysed by—using the software SPSS Statistics (version 16.0) and XLSTAT version 2019.2.1 (XLSTAT, 2019) and a—comparison of means were done at 5 % significance level with Duncan's multiple range test. The principal component analysis (PCA) [9] was performed with seed germination and growth traits as influenced by different PGPR inoculation in cotton.

3. RESULTS AND DISCUSSION

3.1. Effect of PGPR strains on Germination efficiency

Inoculation with dDifferent PGPR strains inoculation—significantly influenced the germination efficiency of cotton seeds. Experimental results in _from—Table 1. revealed that seeds treated with PGPR strains showed significant improvement in germination percentage. Cotton seeds inoculated with strain of PPFM TNAU 1 (T3) showed maximum germination percentage followed by seeds inoculated with Azospirillum sp7 strain (T2) with-i.e., 96.8 and 95.3_%, respectively. Likewise, cotton seeds treated with different PGPR strains positively influenced the mean germination time (day)—(Fig. 1). Mean germination time determines how faster the seed population—germinates in shorter period

the less amount of time [10]. The mean germination time was found to be lower in the strains treated with PPFM TNAU1 (T3) (5.6 days) followed by Azospirillum sp7 (T2) strain (5.9 days) treated seeds whereas seeds ee-inoculated with both Azospirillum sp7 + Phosphobacteria PS1 + Potash bacteria KRB9 (T4) strains resulted in recorded higher mean germination value of 6.3 days. The seeds treated with individual strains of PPFM TNAU1 and Azospirillum sp7 strain had recorded about 16.2 and 14.4% increase, respectively, over the control. Results obtained coincide The present study is in corroboration with those obtained e findings of by Prathibha and Siddalingeshwara. [11]n and -Sirohi et al. [12] where that seed inoculation of Bacillus sp. and Pseudomonas significantly improved germination percentage and the rate of germination in sorghum and wheat seeds. Similar findings were reported by Nehra et al. [13], -El-Sheekh et al. [14], Gowtham et al. [15] and Roman-Ponce et al. [16]



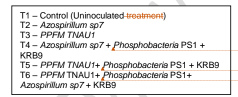


Fig.1. Effect of PGPR strains on mean germination time (MGT)—(days) on cotton seeds.

Inoculation of PGPR strains showed exhibited significant improvement in germination index (GI), germination rate index (GRI) and coefficient of velocity of germination (CVG) (Fig. 2. and Fig. 3.). Cotton seeds inoculated with PPFM TNAU 1 (T3) strain showed maximum germination index (354.3) followed by the treatment T2, Azospirillum sp7 strain (337.3). GI accentuates both germination percentage and speed. Higher GI values indicates a higher germination percentage and rate at which germination occurs [7]. Germination rate index was recorded higher in the treatment T3, PPFM TNAU1 strain followed by the treatment T2 Azospirillum sp7 strain treated seedplants i.e., 17 % day, and 16.2 %-day_1 respectively. The GRI measures the percentage of seeds that germinate on each day of the germination period, with higher GRI values indicateing faster germination [5]. Similarly, the coefficient of velocity of germination also recorded higher in the (T3) PPFM TNAU 1 strain (17.8) inoculated seedsplants. The CVG indicates the speed at which seeds germinate. It increases as the number of germinated seeds increases whileand_the time needed for germination declines [6]. The inoculation of PPFM TNAU1 and Azospirillum sp7 strain had recorded about 27.1 and 21 %, 23.5 and 18.3_%,10.7 and 4.8_% increase ment in germination index, germination rate index and coefficient of velocity of germination over the uninoculated control, respectively. The increased germination efficiency is due to the synthesis of gibbereline, the hormone, gibberellin, which stimulates the activity of alpha-amylase, the germination-specific enzyme ands, alpha-amylase, protease and nuclease which is mainly involved in starch breakdown and assimilation [17], and mitochondrial enzyme activities [18]. The present study coincides is in correboration with results of Xiao et al. [19], and; Hossain et al. [20] in which PGPR inoculation significantly improved germination percentage and germination rate index over the with reference to the control in rice. Similar findings were reported by Makhaye et al. [21], and Hamidi et al. [22], where that seed inoculation of PGPRs significantly improved all-the germination traits such as germination percentage (GP), germination index (GI), germination rate index (GRI), mean germination time (MGT) and coefficient of velocity of germination (CVG) in maize. However, the increased germination traits were also observed in combined inoculants of PGPR strains. The results of the present study are similar to the findings of Anitha-[23] and Meena et al. [24].

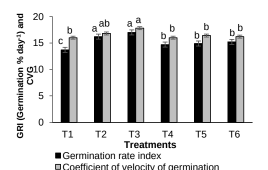
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T1 – Control (Uninoculated treatment) T2 – Azospirillum sp7

T3 – PPFM TNAU1

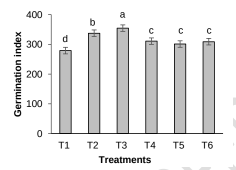
T4 - Azospirillum sp7 + Phosphobacteria PS1 + KRB9

T5 – PPFM TNAU1+ Phosphobacteria PS1 + KRB9

T6 – PPFM TNAU1+ Phosphobacteria PS1+ Azospirillum sp7 + KRB9

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Fig. 2. Effect of PGPR strains on germination rate index GRI (Germination % day.1) and coefficient of velocity of germination of cotton seeds.



T1 - Control (Uninoculated treatment)

T2 – Azospirillum sp7 T3 – PPFM TNAU1

T4 – Azospirillum sp7 + Phosphobacteria PS1 + KRR9

T5 – PPFM TNAU1+ Phosphobacteria PS1 + KRB9

T6 – PPFM TNAU1+ Phosphobacteria PS1+ Azospirillum sp7 + KRB9

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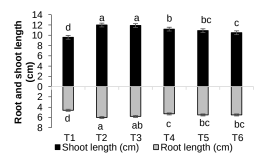
Fig. 3. Effect of PGPR strains on germination index (GI) one cotton seeds.

3.2. Effect of PGPR on seedling emergence traits

Cotton seeds treated with different PGPR strains exhibited significant improvement in the seedling emergence traits such as shoot and root length (Fig. 4.), and seedling vigour index (Table-1). The experimental results obtained indicated that shoot and root length of PGPR inoculated seedlings were significantly higher compared to the uninoculated treated control (Fig. 4). The treatment_rof Azospirillum sp7 strain (T2) inoculated seedsplants resulted in corded-maximum shoot length and root length (12.0 and 6.0 cm, respectively) followed by the treatment, 3 -(PPFM TNAU1) (T3) (with 11.9 and 5.8 cm, respectively). The combined inoculation of PGPR strains (T4, T5 and T6) resulted corded in higher shoot length and root length compared to the non-inoculated control. The percent increment in shoot and root length was 24.4 and 42.8 % in the Azospirillum sp7 (T2) strain inoculated seeds plants over the uninoculated control. PGPR inoculation stimulates the production of the phytehormone IAA, (-Indole-3 Acetic Acid (IAA) and acquiring esmore nutrients such as nitrogen and phosphorus, which, in turn, increases root length and shoot length [25]. In additions of The increasing e in shoot and root length, IAA is mainly due to the primary synthesis of plant hormone, IAA which is responsible for cell elongation in Azospirillum treated plants. The results present findings are similar to those obtained by in accordance with Efthimiadou et al. [26], Taha et al. [27], and El-Gamal et al. [28], Similar findings were reported by Dhale et al. [29], Zamioudis et al. [30], and Pindi et al. [31], where plant growth promoting bacterial (PGPB) inoculation stimulates the production of plant growth hormones that favours root growth and alters root morphology. The plants inoculated with the treatment T3, PPFM TNAU1 strain decumented resulted in higher seedling vigour index (1714.4) as it has higher germination percentage, followed by the treatment T2, Azospirillum sp7 strain (1709.6) which as it showed higher values in shoot and root length compared to other treatments and control. The findings Results obtained by by Noumavo et al. [32] are similar in accordance with the results obtained in the present study. This enhanced seedling vigour index could

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be associated with greater production and metabolism of hormones, auxin and cytokinin which primarily promotes the cell elongation and cell division induced by PGPR inoculation [33,34].



T1 – Control (Uninoculated treatment)

T2 – Azospirillum sp7

T3 - PPFM TNAU1

T4 – Azospirillum sp7 + Phosphobacteria PS1 +

T5 – PPFM TNAU1+ Phosphobacteria PS1 + KRB9 T6 – PPFM TNAU1+ Phosphobacteria PS1+ Azospirillum sp7 + KRB9

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Fig. 4. Effect of PGPR strains on shoot and root length (cm) on cotton seed.

3.3. Effect of PGPR strains on growth traits

Experimental rResults obtained showed-revealed that cotton seeds plants treated with PGPR strains had showed substantial improvedment in leaf area, aerial fresh and dry weight and underground fresh and dry weight (Table-1). The treatment T3, PPFM TNAU1 and the treatment T2, Azospirillum sp7 strains inoculated seedsplants had exhibited a higher foliage size compared to the combined inoculation of strains and control plants. PPFM TNAU1 strain (T3) inoculated seedsplants recorded maximum leaf area (17.4 cm²) followed by the treatment T2, Azospirillum sp7 strain inoculated seedsplants (16.1 cm²). The rate of cell division is higher in the PPFM TNAU1 inoculated seedsplants due to by synthesis of cytokinin, the phytohormone, cytokinin which that contributes to the wider leaf surface area. The inoculation of PPFM TNAU1 and Azospirillum sp7 strain resulted in an increase ment of 57.3 and 45.6_%, respectively, over the control. Our results are similar to those obtained by in correboration with Wang et al. [35], and Namwongsa et al. [36].

Maximum aerial fresh and dry weight were recorded in the treatment T3, PPFM TNAU1 strain (3.25 g seedling⁻¹ and 72.8 mg seedling⁻¹), respectively. Likewise, the maximum underground fresh and dry weight recorded was to be higher in the treatment T2, Azospirillum sp7 strain inoculated plants (0.40 g seedling and 6.85 mg seedling). This increase of aerial fresh and dry weight in the treatment T3, PPFM TNAU1 strain, is due to the higher leaf biomass. Azospirillum sp7 (T2) strain inoculated treatment is found to have maximum underground fresh and dry weight due to increased root length by the synthesis of phytohormones that acts on cell division and cell elongation. The maximum leaf area obtained in the treatment T3, PPFM TNAU1 (T3) inoculated seedlings, contributed to the higher total dry matter production (79.6 mg seedling 1) compared to the (T2) Azospirillum sp7 (77.3 mg seedling 1) strain treated seedlings. The present findings are similar to those obtained by in line with Wang et al. [35] where that increased transcriptional levels of auxin biosynthesis genes enhanced the seedling growth thus improved the biomass. Our results of increased seedling fresh and dry weight were similar in harmony withto those reported by -Etesami and Alikhani. [37], Turan et al. [38], Asari et al. [39], and Egamberdieva et al. [25]. This considerable improvement in seedling growth parameters suggests that selective inoculation of PGPRs could be considered as an effective alternative biofertilizer for promoting cotton seed germination, biomass, and yield.

Table 1. Effect of PGPR strains on germination percentage, seedling emergence and seedling growth traits

Treatments	Germin ation Pperce nt %	Seedling vigour index	Leaf Area (cm²)	Fresh weight (g plant ⁻¹)		Dry weight (mg plant ⁻¹)	
				Aerial weight	Underground weight	Aerial weight	Underground weight
T1 – Control	83.3	1149.5	11.0	2.33	0.22	50.3	5.78
T2 – Azospirillum	95.3	1709.6	16.1	3.19	0.40	70.5	6.85
T3 – <i>PPFM</i>	96.8	1714.4	17.4	3.25	0.38	72.8	6.83
T4 – Azospirillum + PSB + KRB	88.5	1459.4	11.7	2.75	0.30	55.5	6.60

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	T5 – PPFM + PSB							
	+ KRB	88.8	1453.5	11.6	2.65	0.35	55.0	6.63
	T6 – <i>PPFM</i> + PSB+ <i>Azospirillum</i> + KRB	91.3	1453.6	11.3	2.61	0.32	54.0	6.60
	Mean	90.7	1490.0	13.2	2.8	0.30	59.7	6.50
	SEd	_1.06**	34.74**	0.63*	8.28**	5.44	1.74**	0.13**
1	CD(P=0.05)	2.26	74.04	1.33	0.18	0.12	3.72	0.29

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Values are mean of replicate. Vs; values followed by the same letter in each column are not significantly different from each other as determined by DMRT (P = 0.05).

3.4. Principal component analysis (PCA) of seed germination and growth traits in cotton influenced by PGPR strains

The PCA analysis was done using the data of seed germination and growth-related variables such as germination percentage (GP), mean germination time (MGT), germination index (GI), germination rate index (GRI), seedling vigour index (SVI), leaf area (LA), aerial and underground fresh (AFW and UFW) and dry weight (ADW and UDW) obtained from six treatments with four replications (Fig. 5.). The PCA showed revealed that the variables were correlated with the principal component value of 96.60_% (PC1 – 87.57_% and PC2 - 9.03_%). The variables that were obtained from seed treatment of Azospirillum sp7 (T2) and PPFM TNAU1 (T3) strains were located in the right side of the scoring plot, showing positive correlation between the components. However_On the centrary, variables of the control treatment, central (T1)_were located in the bottom left quarter of the scoring plot in PCs (Fig. 5A). In leading plot, The plant growth variables were influenced positively influenced by PGPR strain inoculation, positioned at the right side of the plot (Fig. 5B). PCA results confirmed that PGPR strain inoculation positively influenced the germination traits and growth characters, while mean germination time was least correlated with the inoculation of different PGPR strains. Results from PCA recorded that the treatment Azospirillum sp7 (T2) strain, which is located in the right end of the second quadrant, which detailed that PGPR inoculation improved germination efficiency and seedling growth traits.

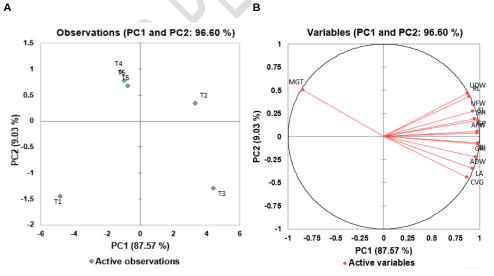


Fig. 5. Principal component analysis (PCA) of cotton as influenced by different PGPR strains (A) Scoring plot of treatments and (B) Loading plot of variables. Variables are GP, Germination percentage; MGT, mean germination time; GI, Germination index; GRI, Germination rate index; SVI, Seedling vigour index; SL, Shoot length; RL, Root length; LA, Leaf area; AFW, Aerial fresh

weight; UFW, Underground fresh weight; ADW, Aerial dry weight and UDW, Underground dry weight.

4. CONCLUSION

Single and combined inoculation of PGPR strain in cotton seeds considerably improved the germination efficiency and plant growth during the early growth stagephase. However, seeds inoculated with individual strain of Azospirillum sp7 and PPFM TNAU1 outperforms the combined inoculation of PGPR strains. These results indicate conclude that the PPGPR strain inoculation has a positive stimulant effect on germination efficiency and plant growth promotion in the cotton crop. It is concluded Thus, the that the investigated beneficial microbes can be employed to increase the physiological potential of cotton to obtain achieve higher yield.

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