In vitro Biofilm Development and Enhanced Rhizosphere Colonization of *Triticum aestivum* by Fluorescent *Pseudomonas* sp.

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Abstract

Effective rhizosphere colonization by plant growth promoting rhizobacteria (PGPR) is a prerequisite for its persistence performance under plant-root soil system. Therefore, to obtain effective strains, we have evaluated twelve strains of fluorescent *Pseudomonas* sp. from wheat rhizosphere exhibiting multifarious plant growth promoting (PGP) activities. These strains formed varying level of biofilm *in vitro* on abiotic surface. Two strains MW3 and MW4 with promising PGP activity but differing in biofilm forming ability. The biofilms were characterized by crystal violet assay, scanning electron and confocal laser scanning microscopy. The MW4 strain, a strong biofilm former is identified by 16S rRNA gene sequence analysis. Strain MW4 showed strong biofilms compared to MW3. Similarly, wheat seeds treated with MW3 and MW4 revealed enhanced rhizospheric colonization by MW4 strain significantly (*P* ≤ 0.05) in higher number (logCFU g⁻¹) up to 45 days after sowing compared to MW3. These findings suggested that strong biofilm forming ability of MW4 might have resulted in enhanced rhizosphere colonization. Therefore, biofilm forming ability of PGPR should be considered as an additional criterion in screening of fluorescent *Pseudomonas* sp.

Keywords: Biofilm; *Pseudomonas fluorescens*; *Triticum aestivum*; root colonization; 16S rRNA gene; confocal laser scanning microscopy.
INTRODUCTION
Rhizobacteria promoting plant growths are diverse and contribute in crop productivity by their various direct and indirect PGP activities. Fluorescent Pseudomonads have been found as one of the important PGPRs exploited in crop production. However, the performance of microbial strains used as biofertilizer is not always consistent due to several biotic and abiotic factors. Therefore, effective rhizosphere colonization by microbial inoculants under field conditions is most critical to promote plant growth consistently. It is now commonly accepted that under natural environment bacteria exist mainly in biofilm mode rather than planktonic state. Biofilms are defined as highly organized microbial communities encased in self-produced extracellular matrix. It is believed that bacterial strains exhibiting biofilm associated characteristics such as flagellar motility, type IV pili, hydrophobicity and production of exopolysaccharides might be helpful in effective biofilm formation and rhizosphere colonization. Therefore, in this study, fluorescent Pseudomonas strains exhibiting multiple plant growth promoting traits were tested for their biofilm formation. Most effective biofilm forming strain (MW4) was further compared with moderate biofilm forming fluorescent Pseudomonas (MW3) to assess possible relationship between biofilm development and wheat rhizosphere colonization.

MATERIAL AND METHODS

Isolation of fluorescent Pseudomonas
Wheat plants cultivated locally in the vicinity of Aligarh city, India were uprooted carefully and rhizospheric soil was collected. Briefly, soil tightly adhered to roots were collected by vortexing the roots in 0.85% NaCl solution to prepare stock suspension. Hundred microlitre of appropriately diluted sample was spread on King’s B agar plates and incubated for 48 h at 28°C. Typical colonies producing yellow-green fluorescence detected under UV light were selected and purified. The cultures were given a strain number and maintained in the laboratory by subculturing. One set of culture was preserved at -80°C in glycerol broth. The fluorescent Pseudomonas strains were biochemically characterized and tentatively identified using standard methods.

Assay for production of plant growth promoting activities
Plant growth promoting activities of isolated bacteria was determined using standard methods as described previously with little modifications. A brief description is given here. Production of Indole-3-acetic acid (IAA) was analyzed using the standard method of Loper and Scarth with the amendment of media by tryptophan (100 µg mL^-1). Siderophore production was studied by growing the cultures in Modii medium. Catechol-type phenolates were measured as described by Reeves et al. Phosphate solubilization was studied using standard method.

The exopolysaccharides (EPS) production was done by the method of Mody et al. 1-aminocyclopropane-1-carboxylate (ACC) deaminase activity of the strains was checked by growing them onto DF (Dworkin and Foster’s) salts minimal agar medium amended with 3.0 mM ACC (Sigma-Aldrich, USA) instead of (NH4)2SO4 as nitrogen source. Bacterial strains were also evaluated for their antifungal activity against Fusarium oxysporum (MTCC-3930), Fusarium ciceri (ITCC-3636), Fusarium solani (ITCC-6953), Aspergillus niger (ITCC-6370), Alternaria alternata (ITCC-6778) by dual culture assay.

Assay for biofilm development
The biofilm formation was determined in 96 well polystyrene microtitre plates as described by O”Toole and Kolter. Briefly, bacteria were grown in the tryptic soy broth (TSB, HiMedia, India) at 28 °C for 24 h. Each well was filled with 200 µL (≈10^7 CFU mL^-1) of culture in fresh medium and allowed to grow for 24 h at 28 °C. The wells were washed with phosphate buffer saline (PBS), followed by staining with 0.1% crystal violet for 20 min. Unbound strain was removed by washing each well with PBS. In each well ethanol (95%) was added to solubilize the bound dye and absorbance was recorded at 590 nm using micro plate reader (ThermoScientific Multiskan EX, UK). Based on absorbance value of dye, the strains were distinguished as strong and moderate biofilm formers according to the criteria adopted by Christensen et al.
Identification of MW4 strain based on 16S rRNA gene sequence analysis

The methods adopted at Macrogen (South Korea) for molecular characterization is briefly described below. Genomic DNA was extracted from MW4 and used as template in a 30 µL reaction mixture by using an EF-TaqDNA polymerase (SolGent, South Korea). The primers 518F (5’-CCAGCAGCGGTATCCTACG-3´) and 800R (5’-TACCCAGGTATCTAATCC-3’) were used for the PCR (Polymerase chain reaction). The PCR reaction was performed with 20 ng of genomic DNA as the template Activation of Taq polymerase at 95°C for 2 min, 35 cycles of 95°C for 1 min, 55°C, and 72°C for 1 min each were performed, finishing with a 10 min step at 72°C. The amplified PCR products were purified with a multi screen filter plate (Millipore Corp., USA). Sequencing reaction was performed using a PRISM BigDye Terminator v3.1 Cycle Sequencing Kit. The DNA samples containing the extension products were added to Hi-Di formamide (Applied Biosystems, CA, USA). Sequencing reaction was performed using a PRISM BigDye Terminator v3.1 Cycle Sequencing Kit. The DNA samples containing the extension products were added to Hi-Di formamide (Applied Biosystems, CA, USA). The mixture was incubated at 95°C for 5 min, followed by 5 min on ice and then analyzed by ABI Prism 3730XL DNA analyzer (Applied Biosystems, Macrogen Inc, Seoul, South Korea). The resulting nucleotide sequence measuring 997 bp in size was submitted to GenBank sequence database (accession number KF553576). The sequence obtained were analysed using Basic Local Alignment Search Tool (BLASTN) and matched against known bacterial sequences present in NCBI GenBank data base (http://www.ncbi.nlm.nih.gov). Related sequences obtained from the database were used to construct a phylogenetic tree using MEGA5 software.

Rhizosphere colonization assay

For colonization study under non sterile soil condition, rifampicin resistant mutant of the two parent strains MW4 and MW3 was selected based on sequential exposure of rifampicin and selecting the mutant on rifampicin amended Kings B agar plate and checked their stability for several generations. *Triticum aestivum* L. (var. PBW 343) surface sterilized seeds were sown in sterile sand-soil mixture for raising seedlings. After 15 days of growth, the plant was uprooted, and roots were washed with sterile distilled water and treated with bacterial culture by dipping the roots in 10 mL diluted culture of fluorescent *Pseudomonas* and MW4 (≈ 10⁵ CFU mL⁻¹) separately and sown in natural loam soil. Collection of rhizosphere and rhizoplane samples (five plants) was made at 15, 30, and 45 days after sowing. Serial dilutions from rhizosphere and rhizoplane samples were prepared separately in normal saline solution (NSS) and plated onto King’s B agar medium containing 800µg mL⁻¹ rifampicin. Untreated plants maintained in similar condition were also subjected to enumerate presence of naturally occurring rifampicin resistant bacteria capable of growing on King’s B agar medium at 800µg mL⁻¹ rifampicin. Only fluorescent colonies grown on rifampicin amended plates were counted.

Biofilm formation on glass cover slips by MW3 and MW4

Crystal violet assay with little modifications as described by O’Toole and Kolter18 was performed to establish the biofilm development on glass coverslips. Briefly, bacterial strains were grown in tryptic soy broth (TSB) as described earlier. A 12 well tissue culture plate with flat bottom containing sterile coverslips were filled with TSB and 100 µL of inoculum (≈10⁷ CFU mL⁻¹) per well and incubated statically at 28 °C for 48 h. After growth coverslips were removed and rinsed in PBS followed by staining with 0.1% crystal violet for 20 min. After staining, coverslips were washed with PBS to remove excess dye and examined under a light microscope (Olympus BX60, Japan) at 100X.

Confocal laser scanning microscopy of biofilms by MW3 and MW4

The biofilms formed on coverslips as mentioned above were stained with acridine orange (Sigma-Aldrich, USA). Microscopic observations and image acquisitions were performed using confocal laser scanning microscopy (CLSM) (Olympus Fluoview FV1000) using standard protocol.

Scanning electron microscopy (SEM) and CLSM and of biofilms formed on the roots of *T. aestivum*

*Triticum aestivum* L. (var. PBW 343) surface sterilized seeds were sown in sterile sand-soil mixture for raising seedlings. After 15 days of growth, the plant was uprooted, and roots were washed with sterile distilled water and treated with bacterial culture by dipping the roots in 10 mL diluted culture of fluorescent *Pseudomonas* and MW4 and MW3 (≈10⁵ CFU mL⁻¹) prepared in TSB. In the control plants, sterile distilled water (SDW) was used instead of culture. After 24 hours the root samples was fixed in 2.5% glutaraldehyde and washed in the same buffer followed by fixation in
osmium tetraoxide. The root were dehydrated by treating with graded concentrations of ethanol and dried in CO₂ with a critical point-dryer, mounted on stubs, and shadowed with gold, (Sputter Coater-Polaron SC7640 UK) for viewing under an scanning electron microscopy (SEM) (Carl ZeissEVO 40, Germany). For CLSM, 24 h-treated root samples were first stained with acridine orange and images were acquired as mentioned above.

**RESULTS**

Isolation, characterization and screening of fluorescent *Pseudomonas*

After initial screening of 56 strains suspected to be fluorescent *Pseudomonas*, twelve strains were selected based on their multiple PGP traits (≥ 3), where MW4 exhibited a comparatively higher level of activity for phosphate solubilization, indole acetic acid, and siderophore production.

![Graphs showing plant growth promoting activities and biofilm formation by fluorescent *Pseudomonas* strains.](image)

Fig. 1. Plant growth promoting activities and biofilm formation by fluorescent *Pseudomonas* strains. (a) Plant growth promoting traits of test strains. (b) Biofilm formation on 96 well microtiter plates under stationary conditions. Bar values followed by a different letters are significantly different (α = 0.05) in Duncan’s multiple range test. MTCC-1749 was used as positive control. (c) Survival and colonization of MW3, MW4 (rifampicin resistant mutant) with wheat rhizoplane and rhizosphere in natural soil conditions. Bar values followed by a different letters are significantly different (α = 0.05) in Duncan’s multiple range test.
(Fig. 1a) along with antifungal activity against *Aspergillus niger*, *Fusarium oxysporum*, and *Fusarium ciceri*. Further, these 12 strains were tested for biofilm forming ability in vitro on polystyrene microtiter plate. The data obtained indicates variations in their biofilm forming ability. The strains were grouped into moderate biofilm former (MW3, MW5, MW6, MW8, MW10, and MW11) and strong biofilm former (MW1, MW2, MW4, MW7, MW9, and MW12) (Fig. 1b).

**Biofilm studies on *Pseudomonas fluorescens* MW4 and MW3 strains in vitro**

On the basis of biofilm forming capability in vitro, plant growth promoting traits and effect on seed germination and seedling growth, two strains (MW4 and MW3) were selected for further study. Biofilm formation by *P. fluorescens* MW4 and MW3 were further performed on glass coverslips (Fig. 2a). The results demonstrated that strain MW4 developed dense and thick biofilms. Further, biofilm was analyzed by CLSM (Fig. 2b), which showed bacterial aggregation and formation of biofilm by MW4 whereas less surface adhered bacteria were observed in MW3 (moderate biofilm former). The average thickness of the MW4 biofilm was found to be 19.29µm as evident from Z-stack analysis (Fig. 2c).

**Root colonization by strong and moderate biofilm forming bacterial strains**

Root colonization was performed by selected two strains MW4 and MW3. The *Pseudomonas fluorescens* MW4 treated seeds yielded log 8.44 CFU g⁻¹ seed. The rhizospheric and rhizoplane viable count of the strain was carried out at 15, 30 and 45 days after sowing and change in bacterial population was observed (Fig. 1c). In rhizoplane, the population of MW4 decreased one log after 15 days of sowing (log 7.38 CFU g⁻¹ root wt) compared to MW3 (log 6.11 CFU g⁻¹ root wt). There was an increase of CFU count after 30 days of sowing but decrease after 45 days. Whereas, in rhizosphere the mean values of viable counts for MW4 and MW3 were log 5.44 and 4.04 CFU g⁻¹ of rhizosphere soil respectively at 15 days after sowing. At 30 days, a little increase in the population of both strains was recorded but a declining trend was observed after 45 days of sowing. Strain MW4 colonized both rhizoplane and rhizosphere more effectively as compared to MW3 even up to 45 days after sowing.

**Biofilm studies of MW4 and MW3 in vivo**

The behavior of the MW4 and MW3 strains during their interactions with wheat seedlings was evaluated using SEM and CLSM. Root sections of wheat bacterized with MW4 and MW3 were examined. The results revealed that cells of MW4 were consistently distributed on the surface of roots (Fig. 3d-j), whereas MW3 showed decreased root colonization and biofilm formation (Fig. 3a-c). A biofilm consisting of bacterial cells and a net-like material suggest extracellular matrix formation (Fig. 3h-j). Less adherent microbial cells were observed in the negative control (untreated) on any of the observed root samples (Fig. 3k-n). Similar pattern of biofilm on root surface was revealed for MW4 when analyzed with CLSM (Fig. 4c-d) compared to MW3 which showed moderate...
biofilm formation (Fig. 4a-b). The average thickness of the MW4 biofilm formed on wheat root surface was found to be 24.92 µm as evident from Z-stack analysis (Fig. 4e). The strain was found a strong biofilm former as well as colonized roots by dense biofilm development compared to moderate biofilm former MW3.

Fig. 3. Scanning electron micrographs of *Triticum aestivum* seedling roots colonized by *Pseudomonas fluorescens* MW3 and MW4. (a) Colonization and biofilm formation on the surface of the root zone by MW3 (moderate biofilm former). (b) Magnified image of the framed region shown in figure 3a. (c) Colonization and biofilm formation of the framed region shown in figure 3b. (d) Colonization and biofilm formation on *Triticum aestivum* roots by MW4. (e) Magnified image of the framed region shown in figure 3d. (f) Large scale colonization of the framed region shown in figure 3e. (g) Biofilm formation by *Pseudomonas fluorescens* MW4. (h-i) Extracellular matrix production by biofilm forming MW4 (white arrow). (j) Magnified image of the framed region shown in figure 3i. (k-n) SEM image of root zone of control (untreated) seedling. (l) Magnified image of the framed region shown in figure 3k.
DISCUSSION

Biofilm forming rhizobacteria are expected to colonize plant roots more stably and can remain active more consistently. To test this hypothesis, fluorescent *Pseudomonas* strains isolated from wheat (*Triticum aestivum* L.) rhizosphere and exhibiting multiple PGP activities and biofilm forming ability *in vitro* were studied for their potential for rhizosphere colonization through biofilm formation.

These strains were tested and identified as *Pseudomonas* sp. and strain MW4 was identified as *Pseudomonas fluorescens* by molecular techniques. The strain MW4 was further selected due to strong biofilm formation ability and increased level of PGP activities.

PGPR including fluorescent *Pseudomonas* are known to exert plant growth promoting properties by both direct mechanisms (assisting nutrient acquirement, adjusting phytohormone...
concentration, nitrogen fixation, phosphate solubilization and sequestering iron) and indirect mechanisms such as production of antibiotics, lytic enzymes and siderophores, competition, induced systemic resistance\(^1\). *Pseudomonas fluorescens* MW4 produced auxins, solubilized phosphate, siderophore and ACC deaminase activity. Presence of multiple PGPR traits observed in this study is in agreement with the report of Zabihi *et al*\(^2\). Significant production of exopolysaccharides by PGPR is expected to confer protection against the strained conditions and assist in bacterial root colonization\(^21\). Moreover, exopolysaccharides have water retaining and adhesive characteristics and play a significant role in stabilization of soil aggregates and the regulation of nutrients and water flow across plant roots\(^22\). Strain MW4 was assessed for its potential to form biofilm on roots of the wheat plant under natural aseptic conditions which is evident from SEM and CLSM. The strain MW4 colonized plant root and form biofilm, indicating that such activity might be helpful in protecting the plant from root borne pathogens\(^23\). To evaluate this, we perform an experiment for their survival in rhizoplane and rhizosphere. The findings have clearly demonstrated better rhizosphere colonization by strong biofilm former MW4 compared to moderate biofilm former, MW3 under natural condition.

**CONCLUSIONS**

Based on above findings it can be concluded that *Pseudomonas fluorescens* MW4 is a strong biofilm former and competent root colonizing PGPR which could be developed as inoculant to assess its performance under field condition. Further mutants of these cultures deficient in biofilm formation need to be tested to explore the role of biofilm and other traits in improving the rhizosphere ecological fitness of the bacteria.

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**CONFLICT OF INTEREST**

The authors declares that there is no conflict of interest.

**AUTHORS’ CONTRIBUTION**

All authors listed have made a substantial, direct and intellectual contribution to the work, and approved it for publication.

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None.

**DATA AVAILABILITY**

All datasets generated or analyzed during this study are included in the manuscript.

**ETHICS STATEMENT**

This article does not contain any studies with human participants or animals performed by any of the authors.

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