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Orchids of Hottest Hot Spot in India: A Rich Source of Endophytes

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Abstract: *Western Ghats is one of the major tropical evergreen-forested regions in India and is one of the hottest hot spots in the World. Increasing urbanization as well as deforestation and thereby resulting habitat loss leads us to give a hand on conservation of such challenging biodiversity in these region. *Aerides crisa* and *Cleisostoma tenuifolium* are two vulnerable orchid plants from Kodagu, which is the heart of Western Ghats and are used as source of endophyte for the study. Isolation and molecular characterization of endophytic bacteria from the plants were studied to its genotypic level by comparing the PCR isolated 16S rRNA sequences. Nucleotide sequences were compared with the NCBI database and accession numbers for each bacterial strain were obtained from the GenBank. About 12 different bacteria were molecularly identified from these two orchids, *A. crisa* and *C. tenuifolium*.*

Orchids and microbes from orchids are less considered areas among researchers. Furthermore, the wide range of endophytes from such vulnerable plants speaks a lot about the necessity of bio-conservation of endophytes, orchids and thereby, harboring habitat of Western Ghats for the upcoming generation. However, vigorous trials were necessary to identify diversified endophytes from vulnerable plant habitats and so as a small step towards the biodiversity conservation of one of the most important biogeographic zones of the World.

Keywords: *Hottest hot spot, Orchid, GenBank, Biodiversity, Endophytes*

I. INTRODUCTION

As per UNESCO World heritage site, it is one of the eight “Hottest hot-spots” of biological diversity in the world. The mountain range is also known as “Sahyadri” begins from the Songadh town of Gujarat, south of Tapti river and covers approximately 1,600km through the states of Maharashtra, Goa, Karnataka, Kerala and Tamilnadu, ending at Marunthuvazhmalai at Swamithope, near Kanyakumari, the southern tip of India.

Kodagu district in Karnataka is located in the central part of the Western Ghats, situated in south India comprises 50% forest and agro-forestry area. Trees of Western Ghats harbor a variety of epiphytic orchids [1]. The forests of Western Ghats are known to be a varietal storehouse of economically important plants. The tropical climate, heavy rainfall from southwest monsoon and favorable soil factors made the area ideal for the rich biodiversity. The central Western Ghats area of Karnataka covers places viz., Kodagu, Hassan, Chikmagalur, Shivamogga, and Uttara Kannada [2].

Orchids are one of the most beautiful creations on the earth, comprising a unique group of plants valued mainly for the ornamental purpose and also for traditional medicine. These are one among the most threatened of all flowering plants. The Orchidaceae is one of the largest plant families with more than 25,000 species globally. These plants are known for their beauty and medicinal property. Although the large population of orchids is confined to their natural habitats, their number is decreasing because of high demand, habitat destruction and indiscriminate collection [3][4][5]. Evidence of plant-associated microorganisms found in the fossilized tissues of stems and leaves has revealed that endophyte-plant associations may have evolved from the time higher plants first appeared on the earth [6]. Endophytes are microbes that live within the plant tissue without causing any noticeable symptoms of disease. Endophytes have been found in all parts of plants including xylem and phloem. The majority of the endophytes have been isolated from trees, but only a few herbaceous plants and shrubs have been examined for the presence of endophytes. *Rhizobium* and other beneficial microbial diversity of three legumes plants that would help as biofertilizers for the crop from Fabaceae family [7].

Previous studies revealed about the endophytic fungal diversity [8] and a combination of endophytic bacteria and fungi [9] from vulnerable epiphytic orchids from Western Ghats. This investigation documents the diversity of residing endophytic bacteria in two plants via, *A. crisa* and *C. tenuifolium* from the heart of Western Ghats.

II. MATERIALS AND METHODS

Aerides crisa and *Cleisostoma tenuifolium* were used as specimens. Somwarpet agro-forestry region (located at 10.42°N 74.73°E latitude) of Kodagu, Karnataka which is the central part of Western ghats was the study area for plant collection. The parts of the plant can be randomly cut off with a disinfected sickle and placed separately in sterile polythene bags to avoid moisture loss. The materials were transported to the laboratory within 24 hours and stored at 4°C until the isolation procedures were completed.

The collected plant parts were thoroughly washed in running tap water to remove dirt and debris. Fresh healthy leaves and roots were selected for endophyte isolation. Epiphytes were removed from the surface by disinfecting the specimen by 70% ethanol for 1 minute, 4% sodium hypochlorite solution for 3 minutes; 70% ethanol for 30s and two rinses in sterilized distilled water. After removing the excess water, the leaf and root were excised into the size of 0.5X0.5 cm with the help of a sterile blade. A total of 50 segments were screened from each and these segments were placed in petri plates nutrient agar with chloramphenicol (150 mg/L). These plates were incubated at 28°C for 24-48 hours and after getting visible bacterial colonies. Each different colony was cultured separately by a streak plate method for obtaining pure culture. These pure cultures were maintained for further experiments. [10] [11].

Genomic DNA of bacterial culture were isolated by phenol-chloroform method according to Mora [12]. 16S rDNA obtained were quantified and amplified using the primers: Forward BSF: 5'GAGTTTGATCCTGGCTCAGG 3'; Reverse BSR: 5' TCATCTGTCCC ACCTTCGGC 3'. The process of PCR was done using the set up; 10XPCRbuffer: 2.5 µl, MgCl₂:2µl, dNTP's mix (1mMeach):5 µl, Primer (10µM): F-0.5µl, R: 0.5µl, *Taq* polymerase (3U/µl): 0.3 µl DNA template (50ng/µl): 4µl. The PCR programme employed was as follows: primary denaturation for 5 minutes at 94°C; 35 cycles of denaturation at 94°C for 30s; annealing for 30 s, and extension at 72°C for 1 min; and a final extension for 10 minutes at 72°C.

Obtained sequences were compared with the known bacterial sequences from the NCBI database and then submitted to GenBank for accession number.

III. RESULTS

The plant specimen, ie, *Aerides crisa* and *Cleisostoma tenuifolium* before collection were shown in Figure 1 and Figure 3 which is present on the tree trunk. Those specimens after collection and before processing were shown in Figure 2 and Figure 4. From the epiphytic orchid plant *Aerides crisa* and *Cleisostoma tenuifolium*, different plants were used for the isolation of endophytic bacteria. A number of endophytes were obtained. We have selected a few for molecular identification. Those selected from *A. crisa* were AR1, AR2, AR3, AR4 and AL2; and from *C. tenuifolium* were BL1, BL2, BL3, BL4, BL5, BR1 and BR2 (names for temporary convenience).



Fig.1: *A crisa* on tree trunk



Fig.3: *A crisa* after collection



Fig. 2: *C. tenuifolium* on its natural habitat



Fig.4: *C. tenuifolium* before processing

The tissue segments were placed on the nutrient agar and kept in an incubator till visible growth appeared. Different colonies were cultured separately and by continuous culturing pure cultures were obtained and maintained aseptically for further research. Figure 5 to Figure 8 showing the streak plates of endophytes from *A. crispa* and Figure 9 to Figure 16 represents the endophytic bacteria from *C. tenuifolium*.



Fig. 5:AR1(*Bacillus pumilus* MT463728)



Fig.6: AR2 (*Bacillus megaterium* MT540506)



Fig.7:AR3 (*Lysinibacillus fusiformis* MT540507)

Fig.8:AL2(*Bacillus cereus* MT540510)

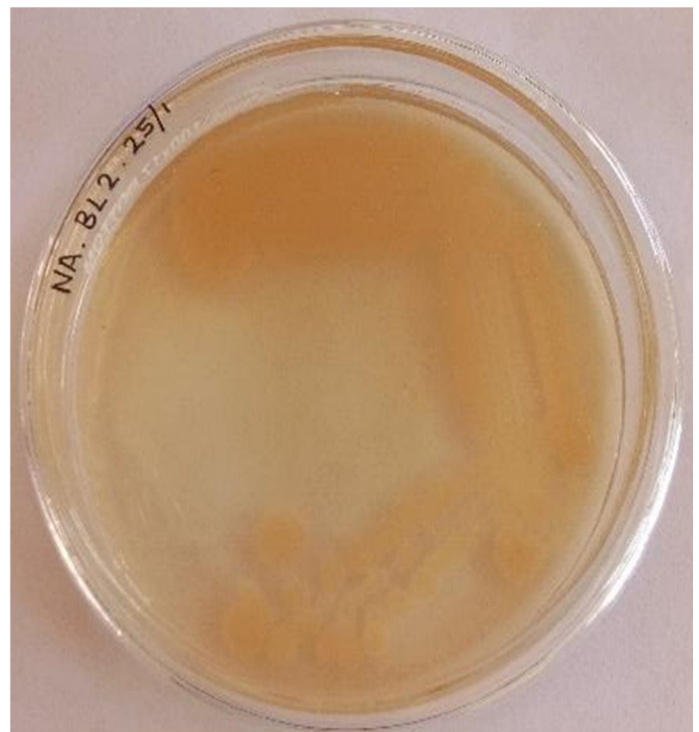


Fig.9:BL1(*Bacillus subtilis* MT463729)

Fig.10:BL2(*Psychrobacillus psychrodurans* MT540515)

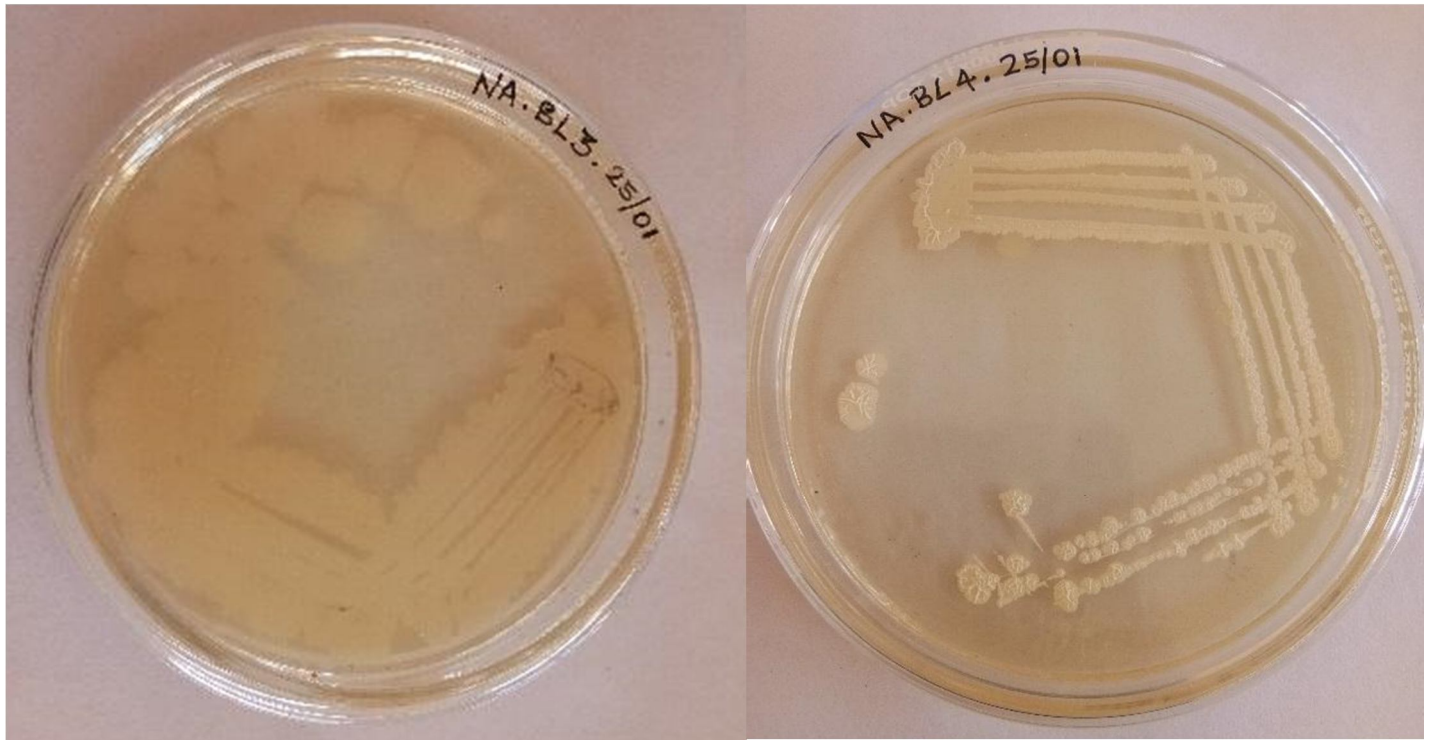


Fig.11:BL3 (*Bacillus cereus* MT540508)

Fig.12: BL4(*Bacillus safensis* MT540512)



Fig.13:BL5 (*Bacillus pumilus* MT540513)

Fig.14:BL6(*Bacillus pumilus* MT463730)

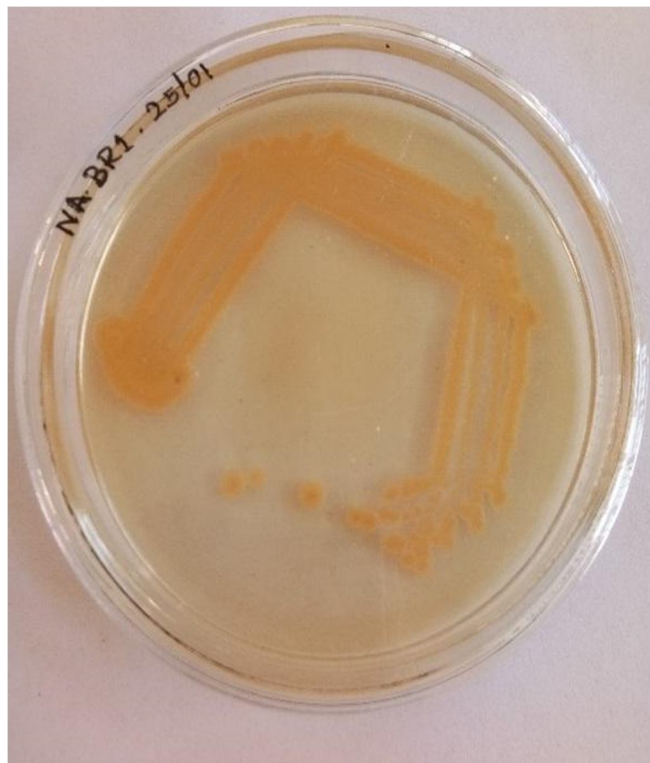


Fig.15:BR1(*Staphylococcus epidermidis* MT463731)

Fig.16:BR2 (*Bacillus pumilus* MT540509)

Genomic DNA of all the bacteria were isolated by phenol-chloroform method. Quantified DNA of five bacteria were amplified using PCR. 16S rDNA were sequenced and the sequences were compared with the known bacterial sequences from the NCBI database. The sequences obtained were submitted to GenBank for Accession numbers; submission code and the accession number of each bacteria from *A.crispa* were given in the Table 1 and from *C tenuifolium* were mentioned in Table 2.

Table 1 showing the GenBank Submission code, Accession number of bacteria from *A.crispa*

Sl. No.	Bacteria	Submission code	Accession number
1.	AR1	SUB7444699	MT463728
2.	AR2	SUB7526079	MT540506
3.	AR3	SUB7526079	MT540507
4.	AL2	SUB7526079	MT540510

Table 2 showing the GenBank Submission code, Accession number of bacteria from *C.tenuifolium*

Sl. No.	Bacteria	Submission code	Accession number
1.	BL1	SUB7444699	MT463729
2.	BL2	SUB7526079	MT540515
3.	BL3	SUB7526079	MT540508
4.	BL4	SUB7526079	MT540512
5.	BL5	SUB7526079	MT540513
6.	BL6	SUB7444699	MT463730
7.	BR1	SUB7444699	MT463731
8.	BR2	SUB7526079	MT540509

From the sequence analysis, twelve strains were studied: four bacterial strains from *A. crispa* were *Bacillus pumilus* MT463728, *Bacillus megaterium* MT540506, *Lysinibacillus fusiformis* MT540507, *Bacillus cereus* MT540510 (Fig.5 to Fig. 8). Eight bacterial endophytes from *C tenuifolium* were *Bacillus subtilis* MT463729, *Psychrobacillus psychrodurans* MT540515, *Bacillus cereus* MT540508, *Bacillus safensis* MT540512, *Bacillus pumilus* MT540513 *Bacillus pumilus* MT463730, *Staphylococcus epidermidis* MT463731, *Bacillus pumilus* MT540509 (Fig.9 to Fig16).

IV. DISCUSSION

Endophytic bacteria are found in roots, stems, leaves, seeds, fruits, tubers, ovules, and also inside legume nodules [7]. Pseudomonadaceae, Enterobacteriaceae, Flavobacteriaceae, Burkholderiaceae, Xanthomonadaceae, and Bacillaceae families are well known plant-associated bacteria [13][14][15][16][17][18]. Joshi and Nongkhaw [19] revealed a definite pattern in the diversity of culturable epiphytic bacteria, host-dependent colonization, microhabitat localization and biofilm formation which play a significant role in plant-microbe interaction. A novel endophytic filamentous bacterium strain was isolated from wild orchid *Grosourda appendiculata* of Thailand [20]. The study of endophytic bacteria is important, not only for understanding their ecological role in their interaction with plants but also for their possible biotechnological applications, such as bioremediation. From this point of view, an interesting interaction between the endophytic bacterial community and glyphosate herbicide was observed during enrichment isolation. Only two bacterial species were recovered from the culture medium supplemented by glyphosate, *Pseudomonas oryzihabitans* and *Burkholderia gladioli*. The bacterium *P. oryzihabitans* was also recovered from total isolation and presented sensibility to glyphosate. This species has been isolated from different samples, such as soil, water, zones of rice cultivation [21], and moreover, soybean seeds [22] [23]. Suggested that these bacterial effects could be potentially useful to promote plant growth during seedling acclimatization in orchid species other than the species of origin [24].

Recently, host specificity has begun to be recognized, using Molecular Plant-Microbe Interactions molecular analysis based on the sequence of ribosomal genes [25] [26] [27]. Extensive information on the molecular mechanisms of other bacteria-plant interactions [28] [29], there is only limited data on the endophyte-host molecular interactions. It would also be interesting to address whether some of the well-studied molecular mechanisms used by phytopathogenic bacteria [30] are to some extent shared with endophytes. Endophytic bacteria provide useful and rich models to study the genetic expression of bacteria in their natural niches or habitats (inside plants), which are more structured and variable than culture media under controlled laboratory conditions. Nevertheless, very little work has been done on this. Genomic projects are being performed on some endophytic bacteria, such as *Azoarcus*.

We have documented about twelve bacterial strains from two epiphytic orchids, *A.crispa* and *C. tenuifolium*. Compared to other methods, the molecular method of bacterial characterization was found better since it compared the genomes, thereby strain differentiation became possible. A vast number of bacterial strains from the small specimen speaks a lot about the importance of biodiversity. The number of studies related to endophytes is very low. Studies related to endophytes and orchids are sparse. Since, the present condition of decreasing forests, followed by reducing orchids leads us to study in this area. Because of the study, we are giving a small hand to biodiversity conservation and thereby conservation of the hottest hot spot, Western Ghats.

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