Characterization of *Bacillus cereus* symbiotic to hemi-parasitic plant
 Santalum album L.

3 Short title: Characterization of *Bacillus cereus* associated with *Santalum album* Linn.

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15 Abstract:

16 **Aims:**

Santalum album L., known as sandalwood plant (white sandal) belongs to the family Santalaceae, is characteristically a hemi-parasite that requires host plant in the early stages for the better growth and development. Besides its extreme economic importance, significant work has not been done to reveal the relationship of beneficial microorganisms with this plant for their better growth and development. Present investigation is an attempt to isolate and characterize the rhizospheric soil bacteria of *Santalum album* L. occurring in some areas of Bankura district of West Bengal, India.

24 Methodology:

The microbial colonies in the soils were estimated as colony forming units (cfu/g dr. soil) from plates prepared by different medium. Phenotypic, biochemical and molecular characters of the bacteria were studied following standard methods. The physico-chemical parameters, and microbial population was determined on the rhizospheric soil of the hemiparasitic sandalwood plant *Santalum album* Linn. occurring at four locations of Bankura district in West Bengal, India.

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32 **Results:**

The population diversity of cultivable heterotrophic, Gram negative, nitrifying, phosphate 33 solubilizing, starch hydrolyzing, spore forming bacteria were higher at Hirbandh with higher 34 organic carbon level than other three locations. Bacterial population was comparatively lower in 35 Basudevpur due to lower water holding capacity. One spore forming bacterium (SW1) was 36 isolated from Hirbandh soil. The isolate (SW1) was characterized by phenotypic properties, 37 scanning electron microscopy, biochemical properties, analysis of fatty acid methyl esters and 38 39 16S rRNA gene sequence and identified as Bacillus cereus (KT626448) which branched with 40 Bacillus cereus BSFN12r (KM405329) with 100% bootstrap support.

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42 **Conclusion:**

It can be concluded that present study is to isolate and characterize the rhizospheric soil bacteria from economically important plant *Santalum album* L. Further studies may find out the positive role of the symbiotic association of *Bacillus cereus* (SW1) with the root of *S. album* as a key factor for the better growth and development of this economically important plant occurring in Bankura district, West Bengal, India.

48 Key words: Santalum album L., Rhizospheric bacteria, Bacillus cereus (SW1), Scanning
49 Electron Microscopy, 16S rRNA gene sequence

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51 Introduction

Santalum album L., commonly known as sandalwood plant (white sandal) belongs to the family
 Santalaceae comprising of more than 15 species and their variants. It is characteristically a hemi-

parasite and requires host plant in the early stages for the better growth and development. Rao 54 (1911) reported that sandal seedlings were incapable to grow beyond one year without haustoria, 55 and confirmed its selectivity for host as although almost all plants in its surroundings may be 56 attacked but the better growth was observed in association with Pongamia pinnata, Albizia 57 lebbeck, Tectona grandis etc (Rama rao et al.1911). Though the plant grows naturally in wide 58 agroclimatic conditions like warm desert of Australia, dry and monsoon climate of India, 59 Vanuatu, eastern Indonesia, subtropical climate of Hawaii and New Caledonia which receives 60 almost uniform rainfall but in India the plant is mostly restricted in southern part only. Presently, 61 there are a few patches in one or two districts of West Bengal have the plant, but it can be 62 introduced in many more areas of West Bengal because it can adapt to various soils although 63 prefers light to medium and well drained soil (Merlin et al., 2006, Rao et al. 2011). Fragrant 64 65 wood and essential oil obtained from sandal are used for the preparation of incense, perfumes, carving and medicine. In spite of its extreme economic importance, significant work has not been 66 done to reveal the relationship of beneficial microorganisms with this plant for their better 67 growth and development. Present study is an attempt to isolate and characterize the rhizospheric 68 69 soil bacteria of Santalum album L. occurring at some areas of Bankura district of West Bengal, India in relation to its role on the growth of the plant. 70

71 Materials and methods

72 Site of soil collection

The rhizospheric soil samples were collected from four different places viz., Hirbundh,
Basudevpur, Bagaldhara and Maitybundh of Bankura district of West Bengal, India.

75 Isolation of the bacteria from soil

100 g of soil samples from the rhizosphere of the plant were collected from different areas of 76 77 Bankura district of West Bengal, India. Soil samples were mixed thoroughly and the soils were put separately in sterile polythene bags, sealed with rubber bands and analyzed in the laboratory. 78 79 Viable aerobic bacterial population were assessed from the plates prepared from the soil suspensions after incubating the plates at 30 ± 0 °C for required days (3-21 d) in a BOD incubator. 80 The microbial colonies in the soils were enumerated as colony forming units (cfu/g dr. soil) from 81 plates prepared with 10 µl soil suspension (10⁻²) mixed with 100 ml of different medium. Soil 82 83 suspension was heated at $60^{\circ}C \pm 0.1^{\circ}C$ for 30 minutes for enrichment culture of the spore

formers. Gram negative bacteria were determined in nutrient agar (NA) (g/l: peptone 5, beef 84 extract 3, agar 2, pH 7.0) and crystal violet (peptone 5 g/l, beef extract 3 g/l, lactose 10 g/l, 85 crystal violet 0.0033 g/l, agar 15 g/l, pH 6.8 \pm 0.1) was added to the medium before plating 86 (Pelczar et al., 1957; Lacey1997; Dangar et al., 2010; Chatterjee et al, 2012; Chatterjee et 87 al,2015). To determine gram-negative bacterial population, crystal violet (0.01 g/l) was added to 88 the medium before plating. The nitrifying bacterial population were assessed on Winogradsky's 89 medium containing $(NH_4)_2SO_4$ (1.0 g/l) and the colonies were visualized (pink colour) by 90 flooding the plates with sulphanillic acid reagent (Pelczar et al., 1957). Nitrifying bacterial 91 colonies were recorded from 5-30 d (5 d intervals) but other colonies were counted after 3 day of 92 incubation. The inorganic phosphate solubilizing bacteria were assessed from the halo zone 93 formation around the bacterial colonies on the insoluble phosphate $[Ca_3 (PO_4)_2]$ containing 94 medium Pelczar et al. (1957). The asymbiotic nitrogen fixing bacterial populations were 95 determined on the nitrogen free medium Pelczar et al. (1957). 96

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98 Characterization of the bacterial isolate

99 The predominant bacterial colonies isolated from the medium were purified and characterized.

The cultural and morphological characters viz. shape, size, elevation, margin, colour, opacity 100 and consistency of the colonies were recorded. Phenotypic and staining properties of the bacteria 101 were studied. Antibiotic sensitivity tests were done with standard antibiotic discs (Brown 2007). 102 103 The bacterial isolate was identified on the basis of biochemical properties, FAME analysis (MIS,MIDI,Sharlock®USA) and 16S rRNA gene sequence analysis (Janssen 1994). The spore 104 105 forming bacterial isolates were observed under scanning electron microscope. The smear preparation of bacterial suspension was done on a cover glass, air dried and heat fixed over a 106 107 flame for one to two seconds followed by 2.5% glutaraldehyde (aquous) for 45 min. Slides were then dehydrated passing through 50%, 70%, 90% and finally with absolute alcohol for 5 min. 108 109 each. Then the gold coated suspensions were observed under scanning electron microscope (HITACHI S-530). For the fatty acid methyl ester analysis (FAME), whole cell fatty acids were 110 111 converted to methyl ester and analysed by gas chromatography. The fatty acid methyl ester 112 composition of bacterial isolates was compared to Sherlock library of known bacterial strains in order to find a closest match. 113

114 **16S rRNA gene sequence analysis**

115 Pure cultured colony of bacterial isolate SW1 was picked up with a sterilized toothpick, suspended in 0.5 ml of sterilized saline in a 1.5 ml centrifuge tube and centrifuged at 10,000 rpm 116 for 10 min. After removal of supernatant, the pellet was suspended in 0.5 ml of Insta Gene 117 Matrix (Bio-Rad, USA), incubated at 56°C for 30 min and then heated 100°C for 10 min. After 118 heating, supernatant was used for PCR. The PCR reaction was prepared with 1µl of template 119 DNA in 20 µl of PCR reaction solution using primers and amplified for 35 cycles at 94°C for 45 120 sec., 55°C for 60 sec. and 72°C for 60 sec. which produced about 1,400 bp DNA fragment. 121 122 dNTPs from PCR products were purified by using Montage PCR clean up kit (Millipore) and the purified PCR products were sequenced using the forward and reverse primers. 123

124 **Results and discussion**

Population dynamics of microorganisms in rhizospheric soils of *S. album* showed that the total aerobic heterotrophic bacteria ranged from 2.9 $\times 10^7$ to 3.3×10^7 cfu/g, the nitrifying bacterial were also higher ranging from 5.2 $\times 10^6$ cfu/g in Basudevpur to 5.7 $\times 10^6$ cfu/g in Hirbandh, phosphate solubilising bacteria ranged from 7.9 $\times 10^2$ cfu/g (Basudevpur) to 8.3 $\times 10^2$ cfu/g (Hirbandh), starch hydrolyzing bacteria ranged from 8.0 $\times 10^4$ to 10.0 $\times 10^4$ cfu/g and spore forming bacteria ranged from 7.5 $\times 10^4$ to 8.0 $\times 10^4$ cfu/g (Table 1).

The organic carbon level varied from 0.37 to 0.51%, nitrogen, phosphorus, potassium and soil pH were also recorded (Table 2). The soil physico-chemical parameters, as well as, the soil types are important factors which influence the soil microbial community. Rhizospheric soil is a hot spot of bacterial diversity and harbours those bacterial strains that may have some impact on soil functional status, as well as, the growth of the plant.

136 The bacterial isolate (SW1) was found in all rhizospheric soil amples of S. album of all locations throughout the year. The bacterial colony was off-white with smooth margin and vegetative cell 137 138 was more than 1 µm and rod shaped. Under SEM study only spores were visualized but no crystal was detected (Fig.1). The isolate was positive for the tests of catalase, methyl red-139 140 Vogues-Proskauer, starch, casein and gelatin hydrolysis and negative for citrate utilization test (Table 3). The isolate reduced nitrate to nitrite which reflects its role nitrogen metabolism in the 141 142 rhizospheric soil (Table 3). The strain was sensitive to nalidixic acid (30 µg/disc), doxycycline (30 µg/disc), bacitracin (10 µg/disc) and tetracycline (30 µg/disc), and resistant to amoxycillin 143

(10 µg/disc), ampicillin (10 µg/disc), polymyxin-B (50 µg/disc) and nystatin (100 µg/disc)(Table 144 3).On the basis of morpho-physiolological and biochemical properties, the bacterium SW1 was 145 identified as Bacillus sp. Through the FAME analysis, 16:1w7c alcohol, 17 isow10c fatty acid 146 supports that the organism belongs to the genus Bacillus and the 17:1 iso w5c, 12:0 iso fatty 147 acid supports that the organism belongs to the species *Bacillus cereus* (Fig. 2). Phylogenetic 148 affiliation of the bacterial isolate reveals that Bacillus sp. SW1 (KT626448) branched with 149 150 Bacillus cereus BSFN12r (KM405329) with 100% bootstrap support (Fig. 3) which confirmed 151 the identity of SW1 as *Bacillus cereus* (Logan et al, 2009). The nucleotide base composition of 16 rRNA gene sequence of the bacterial isolate SW1 was determined which revealed that the AT 152 and GC content were 47.16% and 52.84%, respectively (Table 3). 153

154 It has already been reported that different strains of Bacillus act as plant growth promoters for Saccharum officinerum sugarcane (Dhanraj et al., 2013) and Triticum aestivum (Rawat et al., 155 156 2011) belonging to the family Poaceae. B. cereus has been proved to be a growth promoting rhizobacteria of some plants viz.. Brassica juncea (Aziz et al., 2012), Arabidopsis thaliana (Niu 157 158 et al., 2011), Sophora alopecuroides (Zhao et al., 2011) and Allium ascalonicum (Aziz et al., 2012), belonging to the families Brassicaceae, Fabaceae and Lilliaaceae respectively. Different 159 160 strains of *B. cereus* have been established as facultative mosquito pathogens (Krattiger, 1997; 161 Cooping and Menn, 2001; Wirth et al., 2004; Teng et al., 2005, Chatterjee et al., 2008) which 162 can colonize in mosquito larval guts in relation to the control of Aedes aegypti and A. subpictus 163 larvae. But its growth promoting function on sandalwood plant has not known to date. Present study clearly established the strong association of *B. cereus* SW1 with sandalwood plant 164 Santalum album L. 165

166 **Conclusion:**

Present study is to isolate and characterize the rhizospheric soil bacteria from economically important plant *Santalum album* L. It has already been reported that different strains of *Bacillus* act as plant growth promoters for some economically important plants. Further studies may elucidate the positive role of the symbiotic association of *Bacillus cereus* (SW1) with the root of *S. album* as a key factor for the better growth and development of this economically important plant occurring in Bankura district, West Bengal, India.

174 Table 1. Population density (cfu/g dry soil) of different microbial groups in the rhizosphere

175 of *Santalum album* L occurring at different localities of Bankura district

Sl.no.	Types of organisms	Hirbandh	Basudevpur	Bagaldhara	Maity bandh
1.	Aerobic heterotrophic	3.3±0.001	2.9±0.001	3.1±0.001	3.0±0.001
	bacteria(10 ⁷)				
2.	Gram (–) bacteria (10^6)	3.9±0.001	3.5±0.002	3.7±0.001	3.3±0.001
3.	Nitrifying bacteria (10^6))	5.7±0.003	5.5±0.013	5.2±0.002	5.5±0.013
4.	Phosphate solubilising bacteria (10 ²)	8.3±0.014	7.9±0.018	8.1±0.011	8.0±0.020
5.	Starch hydrolyzing bacteria (10 ⁴)	10±0.02	8±0.02	9.1±0.018	8.5±0.021
6.	Spore forming bacteria (10^4)	8±0.012	7.5±0.015	7.8±0.008	7.6±0.018
7.	Fungi(10 ⁴)	4.66±0.002	4.1±0.003	4.5±0.001	4.2±0.011

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177 Table 2. Physicochemical properties of rhizospheric soil

Sl.	Area	Ν	Р	K	pН	Organic	Texture
no.		(kg/Acre)	(kg/Acre)	(kg/Acre)		carbon(%)	
1	Hirbundh	94.67±	30±0.03	6.9±0.013	6.5±0.019	0.44 ± 0.001	Lateritic,
		0.89					hard
							rocky
2	Basudevpur	72.43±0.40	25±0.01	6.35±0.011	6.2 ± 0.001	0.37 ± 0.001	Lateritic,
							hard
							rocky
3	Bagaldhara	82±0.71	50±0.06	7.5±0.02	6.4±0.017	0.51±0.002	Lateritic,
							hard
							rocky
4	Maity	76 ± 0.46	40±0.04	7.1±0.017	6.3±0.013	0.47 ± 0.001	Lateritic,
	bundh						hard
							rocky

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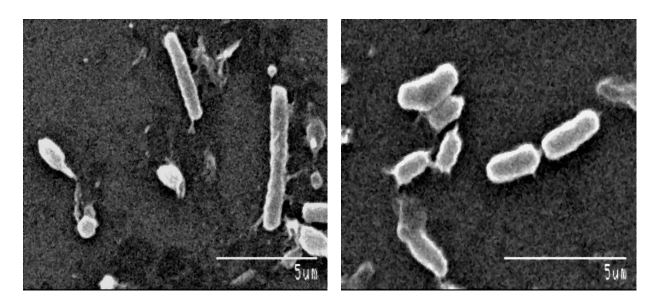
179

Table 3. Biochemical properties of the bacterial isolate (SW1)

Name of the tests	Observations/ Result		

Off white, spherical, elevated			
rod shaped, >1 µm, motile			
- ve			
+ ve			
R			
R			
R			
R			
S			
S			
S			
S			

181 Where, S = sensitive; R = Resistant



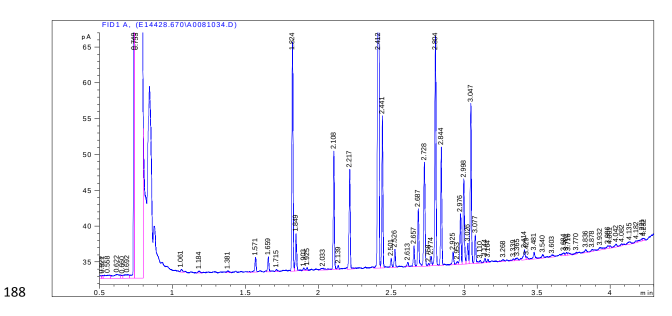


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Plate 1

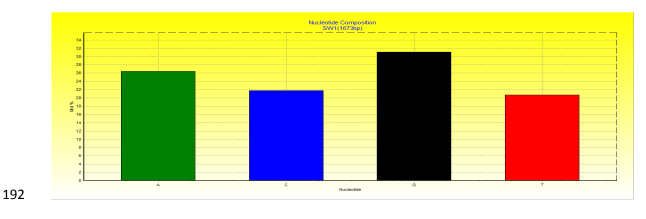


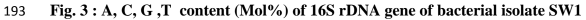
187 Fig. 1: Scanning electron micrograph of *Bacillus cereus* SW1 (Plate 1 and 2)

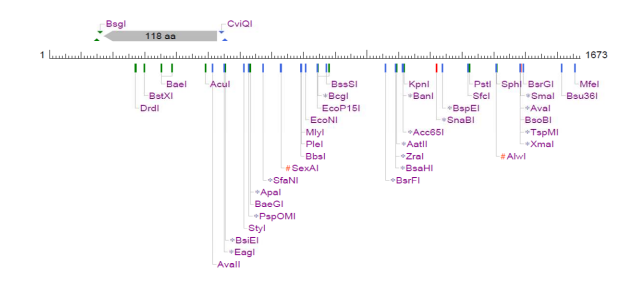


189 Fig 2. FAME analysis of the bacterial isolate SW1

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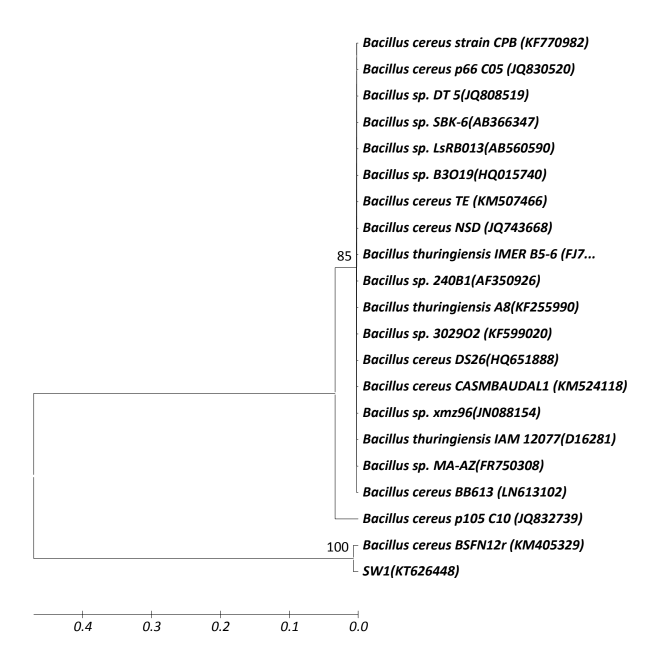


Fig. 5 : Neighbor-joining tree based on 16S rRNA genes sequences of *Bacillus cereus*SW1 (KT626448) strain along with few other 16S rRNA genes retrieved from NCBI.

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