Bacterial endophytes of the medicinal herb *Hygrophila spinosa* T. Anders and their antimicrobial activity

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ABSTRACT

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Aims: The ethnobotanical herb *Hygrophila spinosa* T. Anders (Acanthaceae) is native to India and used in traditional ayurvedic medicines for its pharmacologically important phytochemicals. This study aims to isolate and characterize the culturable bacterial endophytes of *H. spinosa* and evaluate their antimicrobial properties.

Place and Duration of Study: The experiments were performed in the Department of Botany, Serampore College, Serampore as well as in the Microbiology Laboratory, Department of Botany, University of Calcutta, Kolkata during 2011 to 2012.

Methodology: Bacterial endophytes were isolated from healthy plant tissues following surface sterilization and plating on nutrient agar, glycerol asparagine agar and tryptic soy agar. They were characterized physio-biochemically following standard microbiological and biochemical methods. The endophytes were screened for production of antimicrobial compounds following cross-streak assay against test strains *Bacillus subtilis*, *B. cereus*, *Escherichia coli*, *Pseudomonas cepacia*, *Klebsiella pneumoniae* and *Staphylococcus aureus* on nutrient agar plates.

Results: Eleven phenotypically distinguishable bacterial endophytes were isolated from surface sterilized leaf, stem and root tissues and Shannon Weaver diversity index clearly revealed more diverse (0.83) types of endophytes in leaves than in stem (0.48) and root (0.41) tissues. Physio-biochemical features of the isolates clearly indicated distinct variation in their sugar fermentation profiles along with NaCl tolerance. The endophytes produced important enzymes like catalase, amylase, gelatinase, nitrate reductase and lipase. The bacterial isolates belonged to the genera *Bacillus, Paenibacillus, Pseudomonas, Ralstonia, Staphylococcus, Micrococcus* and *Acidomonas*. Antibiotic sensitivity profile, however, have indicated that the isolates were mostly resistant to amoxycillin and bacitracin, while they were highly susceptible to tetracycline followed by neomycin and streptomycin. Interestingly, the bacterial endophytes of *H. spinosa* give a definite stamp on their antimicrobial activity against *E. coli* and *K. pneumoniae* followed by *S. aureus*. Two isolates, *Paenibacillus* HGS 202 and *Acidomonas* HGR 302 obtained from stem and root segments respectively showed antimicrobial activity against *B. subtilis, B. cereus, E. coli, K. pneumoniae* and *S. aureus*.

Conclusion: This study identified eleven bacterial endophytes harboring the leaves, stem and root of *H. spinosa* which demonstrated antibacterial activity against Grampositive as well as Gram-negative bacterial strains. Moreover these endophytic bacterial isolates could be exploited as sources of antibacterial substances.

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19 Keywords: Hygrophila spinosa, Endophytic bacteria, Antibacterial activity, Antibiotic 20 sensitivity, Enzyme profile, NaCl tolerance

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21 **1. INTRODUCTION**

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23 Medicinal plants provide valuable therapeutic agents in traditional medicines which are 24 used on a global level for helping with a wide variety of human health issues. Hygrophila 25 spinosa T. Anders, belonging to the family Acanthaceae, is a promising medicinal herb 26 mentioned in ancient ayurvedic literature as having great economic potential. The plant 27 is indigenous to the Indian subcontinent and is reported to contain phytosterols, fatty acids, polyphenols, proanthocyanins, alkaloids, flavonoids, terpenoids, vitamins, and 28 29 glycosides as major chemical constituents. In traditional medicine, H. spinosa is used 30 mainly for the treatment of hyperdipsia, vesical calculi, flatulence, diarrhea, dysentery, 31 leukorrhea, gonorrhea, asthma, blood diseases, gastric problems, cancer, rheumatism, 32 etc. Many essential phytochemicals isolated from the whole plant including lupeol, 33 stigmasterol, apigenin-7-O-glucuronide, apigenin-7-O-glucoside, betulin, 25-oxohentriacontanyl acetate, methyl 8-n-hexyltetracosanoate, oleic acid, linoleic acid, etc. 34 35 have exhibited antitumor, antibacterial, antidiabetic, antiinflamatory, antipyretic, 36 antioxidant and hepatoprotective activity [1, 2].

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38 It has been rationalized that plants having an ethnobotanical history and exploited for 39 human use in traditional medicine may harbor an endophytic population which may 40 produce a plethora of microbial metabolites related closely to the plant biochemistry [3]. 41 Endophytes, by definition, are microorganisms colonizing living internal tissues of plant 42 either symbiotically or in mutualistic relationship. They occur ubiquitously in all plant 43 species on earth and benefit the host plant growth by fixation of atmospheric nitrogen, 44 production of growth promoting substances, imparting effective disease management, 45 plant protection and stress tolerance [4]. In addition recent studies have established that 46 secondary metabolites elaborated by these microbial endophytes could serve as 47 prospective resources of antimicrobial substances, antioxidants, cytotoxic compounds, 48 growth hormones and hydrolytic enzymes of biotechnological applications [5, 6].

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50 In view of the increasing prevalence of antibiotic-resistant human and plant pathogens, 51 there is an escalating demand for newer antimicrobials from natural sources. Bacterial 52 and fungal endophytes residing inside the healthy plant tissues are believed to carry out 53 a resistance mechanism to overcome pathogenic attack and have emerged as a promising source of newer antimicrobial compounds. Several antimicrobial metabolites 54 55 belonging to structural classes like alkaloids, peptides, benzopyranones, flavonoids, 56 phenolic acids, guinones, steroids, terpenoids, tetralones, xanthones, and others have 57 been obtained from endophytes. The occurrence of endophytic bacteria in agricultural or 58 medicinal plants has been reported quite extensively [7-9]. A comparison of different 59 endophytic hosts shows that nearly 35% of the endophytes possessing antimicrobial 60 activity have been isolated from medicinal plants followed by 29% from agricultural crops 61 [6]. The diversity and ecological distribution of fungal endophytes associated with 62 different medicinal plants native to China, Malaysia, Australia and India have been 63 investigated with special emphasis on their antimicrobial efficacy. A mass of bioactive 64 natural products isolated from endophytes have been reported in recent years and 65 majority of them have been derived from endophytic fungi [3, 7, 10, 11]. However, little 66 information is available on the occurrence as well as on the potential significance of 67 bacterial endophytes from medicinal plants. Although, medicinal properties of H. spinosa 68 have been studied in details by many researchers [1, 2], reports on the endophytic 69 population of this medicinal herb is lacking. Biodiversity of both culturable and 70 unculturable endophytic microbial communities of H. spinosa, therefore, needs to be 71 determined. However, culturable endophytic bacterial isolates deserve special attention 72 for further development of microbial-based biotechnological products and formulations. 73 In the present study, we focused on the isolation, characterization and antimicrobial 74 evaluation of bacterial endophytes from H. spinosa.

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78 2. MATERIAL AND METHODS

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80 2.1 Collection of plant samples

Healthy plants of *Hygrophila spinosa* T. Anders (Acanthaceae) were collected from Medicinal Plant Garden of Serampore College, Hooghly, West Bengal and Department of Botany, University of Calcutta, Kolkata in sterile zip lock polythene bags. The collected plants were brought immediately to the laboratory and stored at 4°C until used for the isolation of bacterial endophytes.

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87 2.2 Isolation and characterization of endophytes

88 Fresh and healthy leaf, stem and root segments were cut from the collected plants, washed thoroughly under running tap water. Surface sterilization was performed in 89 sterile glass bottles by consecutive immersion in 70% ethanol (2 – 3 min), 0.5 % sodium hypochlorite (5 -10 min) and again in 70% ethanol for 30 sec [7]. This was followed by 90 91 92 repeated washing of plant samples in sterile distilled water for at least three times. 93 Samples were blot dried on sterile towels and cut aseptically into small sections before 94 plating on previously prepared nutrient agar, glycerol asparagine agar and tryptic soy agar plates for isolation of bacteria. The plates were incubated at 30°C for 2 - 4 days 95 96 and observed for growth of bacterial colonies surrounding the leaf, stem and root 97 sections. Pure cultures of bacterial endophytes were developed by dilution-streaking on 98 the same media and maintained on slopes of nutrient agar for further study. Bacterial 99 strains were characterized and identified following micromorphological and physio-100 biochemical analysis following standard protocols [12, 13].

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102 **2.3 Diversity of endophytes**

103 Based on the total number of samples plated and the number of samples yielding 104 isolates, colonization frequency and isolation rate were calculated. Colonization 105 frequency was calculated as the total number of plant samples infected by bacteria 106 divided by the total number of samples incubated. Isolation rate was determined as the 107 number of bacterial isolates obtained from plant samples divided by the total number of 108 samples incubated. The Shannon Weaver biodiversity index H^{\prime} was calculated as:

109 $H' = -\Sigma Pi X \ln Pi$, where, *Pi* is the proportion of individuals that species "*i*" contributes to 110 the total [7, 14].

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112 **2.4 Antibiotic susceptibility spectrum**

Antibiotic sensitivity test was performed following the Kirby Bauer disc-diffusion assay
method [15] using antibiotic impregnated discs (6 mm diameter) from Himedia (India).
Based on the diameter of inhibition zone recorded to nearest mm, the organisms were
categorized as resistant, intermediate and sensitive following DIFCO Manual 10th edition
(1984). Antibiotics used include: amoxycillin (30 µg/disc), bacitracin (10 U/disc),
chloramphenicol (30 µg/disc), neomycin (30 µg/disc), streptomycin (30 µg/disc) and
tetracycline (30 µg/disc).

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121 **2.5 Production of antimicrobial substances**

122 Bacterial endophytes were primarily screened for production of antimicrobial substances 123 following cross-streak assay method using six test organisms like Bacillus subtilis, B. 124 cereus, Escherichia coli, Pseudomonas cepacia, Klebsiella pneumonia and 125 Staphylococcus aureus [16]. Nutrient agar plates were inoculated with bacterial 126 endophytes as a single streak at the centre of the Petri plate and incubated for 5 days at 127 30°C. Overnight grown cultures of the test organisms were streaked at right angle to the 128 producer endophyte and observed for its growth / inhibition after 24 - 48 h of incubation 129 at 30°C. The length of inhibition zone was measured to nearest mm.

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133 3. RESULTS

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135 **3.1 Diversity of bacterial endophytes**

136 Segments of surface sterilized leaf, stem and root of Hygrophila spinosa (Acanthaceae) incubated on nutrient agar, glycerol asparagine agar and tryptic soy agar plates showed 137 138 growth of morphologically distinguishable bacterial colonies surrounding the segments after 48-96 h. Avoiding the repetitive strains a total of 11 phenotypically distinguishable 139 bacterial endophytes were isolated in pure form from 118 segments (39 leaf, 39 stem 140 141 and 40 root) of *H. spinosa*. Out of these 11 isolates, six were derived from leaf, while 142 stem and root segments yielded three and two isolates respectively (Table 1). The 143 colonization frequency was lower in leaf samples (17.9%) as compared to the stem 144 (20.5%) and root (22.5%), while the isolation rate was poor in root (0.05) but increased gradually in stem (0.07) and leaf (0.15) samples. The Shannon-Weaver diversity index 145 146 showed that leaves (0.83) of *H. spinosa* harbor more diverse types of endophytic 147 bacteria than in its stem (0.48) and root (0.41).

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Table 1. Diversity of endophytic bacterial isolates in leaf, stem and root tissues of Hygrophila spinosa

Parameters		Total		
Farameters	Leaf	Stem	Root	Total
Number of samples	39	39	40	118
Number of sample yielding isolates	07	08	09	24
Number of isolates	06	03	02	11
Colonization Frequency, % ^a	17.9	20.5	22.5	20.3
Isolation Rate ^b	0.15	0.07	0.05	0.09
Shannon-Weaver Diversity Index ^c	0.83	0.48	0.41	0.68

^a Colonization frequency was calculated as the total number of plant samples infected by bacteria divided by the total number of samples incubated. ^b Isolation rate was calculated as the number of bacterial isolates obtained from plant samples divided by the total number of samples incubated. ^c Shannon Weaver diversity index H^{-/} was calculated as: H^{-/} = - Σ Pi X In Pi, where, Pi is the proportion of individuals that species "i" contributes to the total [7, 14].

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Table 2. Micromorphological characteristics of bacteria isolated from leaf, stem and root tissues of Hygrophila spinosa

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Tissue	Isolate	Cell	Gram	Motility	Size, µm	Endospore	Diffusible
	no.	morphology	nature				pigments
Leaf	HGL 101	cocci, in cluster	<mark>positive</mark>	non-motile	0.5 dia	<mark>absent</mark>	none
	HGL 102	cocci, single	<mark>positive</mark>	non-motile	0.4 dia	<mark>absent</mark>	yellow
	HGL 103	short rod	negative	<mark>motile</mark>	0.4 X 0.3	<mark>absent</mark>	green
	HGL 104	rod, single	<mark>positive</mark>	motile	1.1 X 0.3	<mark>present</mark>	none
	HGL 105	short rod	positive	non-motile	0.5 X 0.4	present	none
	HGL 106	short rod	negative	<mark>motile</mark>	0.5 X 0.3	absent	none
Stem	HGS 201	rod, in chain	<mark>positive</mark>	motile	1.1 X 0.5	present	none
	HGS 202	rod, single	<mark>positive</mark>	motile	0.8 X 0.4	<mark>present</mark>	none
	HGS 203	cocci, single	positive	non-motile	0.5 dia	<mark>absent</mark>	yellow
Root	HGR 301	short rod	negative	motile	0.5 X 0.4	absent	none
	HGR 302	short rod	negative	non-motile	0.5 X 0.4	absent	none

163 Colony morphology was detected in Tryptic soy agar medium after 5 days of growth at 32°C.

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164 **3.2 Characterization and identification of isolates**

The bacterial endophytes of *H. spinosa* were characterized 165 based on micromorphological (Table 2) and physio-biochemical characters (Table 3). Out of 11 166 isolates seven were Gram-positive (three cocci and four rod) and four were Gram-167 168 negative (all rod). Filamentous forms were not detected in any of the plant samples. Six 169 isolates out of 11 showed motility and only three produced yellowish to green diffusible 170 pigments during growth on tryptic soy agar plates. All Gram-positive rods showed 171 endospore formation.

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173 Enzymatic profile of endophytic bacterial isolates showed that all of them produced catalase, while about 55 and 64% of the isolates produced amylase and gelatinase 174 175 respectively (Table 3). Lipolytic (55%) and nitrate reductase (36%) activities were not uncommon amongst the endophytic isolates. Production of indole by the enzyme 176 177 tryptophanase was evident only in isolates HGL 103, HGL 105 and HGR 301. The 178 isolates showed wide degree of tolerance to NaCl (2.5 - 10%) in the growth medium. 179 The endophytes were also screened for their ability to utilize and ferment dextrose, 180 fructose, maltose, sucrose and lactose in phenol red agar medium supplemented with 181 1% sugar (Table 4). While dextrose was the best carbohydrate utilized by all most all the 182 bacterial endophytes, lactose was fermented by only two isolates. The endophytic 183 isolates were moderate in fermenting fructose, sucrose and maltose.

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186Table 3.Biochemical characterization of bacterial endophytes from leaf, stem187and root tissues of *H. spinosa*

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Plant tissue	laslata		E	la de la	NaCl			
	lsolate no.	Catalase	Amylase	Gelatinase	Lipase	NO ₃ Reductase	Indole production	tolerance %
Leaf	HGL 101	+	+	+	+	-	-	10.0
	HGL 102	+	-	+	-	-	-	10.0
	HGL 103	+	-	+	+	-	+	3.5
	HGL 104	+	-	+	-	+	-	4.0
	HGL 105	+	-	-	+	+	+	4.0
	HGL 106	+	-	-	-	-	-	4.5
Stem	HGS 201	+	+	-	+	-	-	4.0
	HGS 202	+	+	+	-	-	-	4.0
	HGS 203	+	+	+	-	+	-	10.0
Root	HGR 301	+	+	+	+	-	+	3.0
	HGR 302	+	+	-	+	+	-	2.5

189 *"+" indicate positive response, "-" indicate negative response*

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Table 4. Fermentation of sugars by bacterial endophytes isolated from leaf, stem and root tissues of *H. spinosa*

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Plant	la elete e e	Fermentation of sugars							
tissue	Isolate no.	Dextrose	Fructose	Lactose	Maltose	Sucrose			
Leaf	HGL 101	+	+	-	+	+			
	HGL 102	+	+	-	-	-			
	HGL 103	+	-	-	-	-			
	HGL 104	+	+	-	-	+			
	HGL 105	+	+	-	+	+			
	HGL 106	-	-	+	-	-			
Stem	HGS 201	+	-	-	-	-			
	HGS 202	+	+	-	-	+			
	HGS 203	+	+	+	+	+			
Root	HGR 301	+	+	-	+	-			
	HGR 302	+	-	-	-	-			

195 "+" indicate positive response, "-" indicate negative response

196 Fermentation of sugars was screened in phenol red agar medium supplemented with 1% sugar.

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Based on microscopic and biochemical analysis, the bacterial isolates were tentatively identified as species of *Bacillus* (HGL 104, HGS 201),
 Paenibacillus (HGL 105, HGS 202), *Pseudomonas* (HGL 103, HGR 301), *Ralstonia* (HGL 106), *Staphylococcus* (HGL 101), *Micrococcus* (HGL

- 199 **102, HGS 203) and** *Acidomonas* **(HGR 302)**.
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201 **3.3 Antibiotic sensitivity profile**

Antibiotic sensitivity pattern of the endophytic bacterial isolates was determined by disc-diffusion method against six different antibiotics like amoxycillin, bacitracin, chloramphenicol, neomycin, streptomycin and tetracycline. Results as shown in Table 5 depict that, bacterial endophytes from leaf, stem and root tissues of *H. spinosa* were mostly resistant to amoxycillin and bacitracin, while they were mostly sensitive to tetracycline followed by neomycin and streptomycin. One leaf endophyte, *Staphylococcus* HGL 101 was highly resistant to five antibiotics and was followed by *Micrococcus* HGS 203 showing resistance to four of the six tested antibiotics. On the contrary, the isolates from leaf and stem (*Paenibacillus* HGL 105, *Bacillus* HGS 201 and *Paenibacillus* HGS 202) showed sensitive to intermediate response towards all the tested antibiotics.

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210 Table 5. Screening of bacterial endophytes from Hygrophila spinosa for their antibiotic susceptibility following disc-diffusion assay

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Plant		Diameter of inhibition zone, mm Antibiotics							
	Isolate								
tissue		Amoxycillin	Bacitracin	Chloramphenicol	Neomycin	Streptomycin	Tetracycline		
Leaf	Staphylococcus HGL 101	08 (R)	NIL (R)	9.5 (R)	12 (R)	11 (R)	40 (S)		
	Micrococcus HGL 102	14 (I)	12 (I)	22 (S)	20 (S)	32 (S)	10 (R)		
	Pseudomonas HGL 103	22 (S)	14 (S)	NIL (R)	22 (S)	32 (S)	NIL (R)		
	Bacillus HGL 104	23 (S)	NIL (R)	26 (S)	18 (S)	18 (I)	19 (S)		
	Paenibacillus HGL 105	14 (I)	12 (I)	18 (S)	24 (S)	30 (S)	26 (S)		
	Ralstonia HGL 106	11 (R)	12 (I)	18 (S)	28 (S)	36 (S)	44 (S)		
Stem	Bacillus HGS 201	25 (S)	13 (S)	14 (I)	20 (S)	27 (S)	24 (S)		
	Paenibacillus HGS 202	20 (S)	16 (S)	17 (I)	16 (I)	32 (S)	20 (S)		
	Micrococcus HGS 203	09 (R)	NIL (R)	9.5 (R)	21 (S)	NIL (R)	19 (S)		
Root	Pseudomonas HGR 301	7.5 (R)	NIL (R)	26 (S)	14 (I)	NIL (R)	20 (S)		
	Acidomonas HGR 302	11 (R)	08 (R)	21 (S)	16 (I)	25 (S)	22 (S)		

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213 R=Resistant, I=Intermediate, S=Sensitive

214 Antibiotic susceptibility was tested on nutrient agar plates using antibiotic impregnated discs (6 mm) from HIMEDIA, India

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215 **3.4 Evaluation of antimicrobial activity**

Antimicrobial activity of all eleven bacterial endophytes were assessed against six bacterial test organisms, *B. subtilis*, *B. cereus*, *E. coli*, *P. cepacia*, *K. pneumoniae* and *S. aureus* following cross-streak method on nutrient agar plates. The isolate which inhibited growth of any of the test isolate(s) was considered having antibacterial activity and the length of inhibition zone was measured (Table 6). Out of 11 endophytes screened, majority showed antibacterial activity against *E. coli* and *K. pneumoniae* followed by *S. aureus*. Isolates *Paenibacillus* HGS 202 and *Acidomonas* HGR 302 obtained from stem and root tissues respectively showed comparatively broad spectrum of antibacterial activity inhibiting both Gram-positive and Gram-negative test organisms.

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Table 6. Evaluation of antimicrobial activity of bacterial endophytes of *Hygrophila spinosa* following cross-streak method

		Length of inhibition zone, mm Test organisms							
Plant	Isolate								
tissue	Isolale	Bacillus subtilis	Bacillus cereus	Pseudomonas cepacia	Escherichia coli	Klebsiella pneumoniae	Staphylococcus aureus		
Leaf	Staphylococcus HGL 101	-	-	-	20	10	-		
	Micrococcus HGL 102	-	-	-	-	-	-		
	Pseudomonas HGL 103	-	-	-	-	-	-		
	Bacillus HGL 104	-	-	-	20	10	-		
	Paenibacillus HGL 105	-	-	5	-	-	5		
	Ralstonia HGL 106	-	-	-	-	5	-		
Stem	Bacillus HGS 201	-	-	-	20	20	-		
	Paenibacillus HGS 202	1	1	3	6	-	3		
	Micrococcus HGS 203	-	-	-	20	8.5	8		
Root	Pseudomonas HGR 301	-	-	-	20	5	-		
	Acidomonas HGR 302	4	2	-	20	5	3		

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"-" means no inhibition zone produced

229 DISCUSSION

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231 Studies on the diversity of culturable microbial endophytes in medicinal and vegetative crop 232 plants are essential to understand their potentials and importance in different fields of 233 biotechnology. This study is the first attempt to isolate microbial endophytes from the 234 traditional medicinal herb H. spinosa. We have screened only the medicinally important plant 235 organs like root, stem and leaf of H. spinosa, although endophytes could also harbor in 236 flower, fruit and seeds. The leaves of *H. spinosa* were found to harbor more diverse types of 237 bacterial endophytes than stem or root segments (Table 1). Such species richness in leaves 238 may be attributed to the anatomical peculiarities of the leaves and micro-environmental 239 conditions rich in essential nutrients which drives the selective force for survival of tissue 240 specific endophytic taxa. Similar prevalence of endophytes in leaf tissues have been 241 observed in Paederia foetida [17], Kigelia pinnata [18] and Quercus ilex [19].

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243 Spatial distribution of endophytic genera also depends on seasonal variation, precipitation, 244 soil parameters and location of plants, plant age and genotypes [4]. Here, we have tested 245 only one genotype from cultivated soil of two different localities which does not reflect the 246 true portrait of culturable endophyte diversity of H. spinosa. The phenotypically 247 distinguishable bacterial endophytes harboring leaves, stem and root tissues of H. spinosa 248 were characterized in details (Table 2 - 4) and tentatively identified as members belonging to 249 bacterial genera Bacillus, Paenibacillus, Pseudomonas, Ralstonia, Staphylococcus, 250 Micrococcus and Acidomonas. These isolates were mostly the fast growing endophytes and 251 were also reported to colonize several other host plants. Occurrences of similar endophytic 252 bacterial genera have been reported from medicinal plants like Gynura procumbens, 253 Azadirachta indica, Boerhaavia diffusa, Phyllanthus emblica, P. foetida etc. [17, 20-22]. In 254 addition, several authors have reported the presence of endophytic actinobacteria inside 255 medicinal plants belonging to the genera Streptomyces, Pseudonocardia. 256 Promicromonospora, etc. [23, 24]. However, such filamentous forms have not been recorded 257 during the present study.

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259 Information regarding production of enzymes by microbes of plant origin are few although 260 endophytic bacteria isolated from leaves of maize [25], leaves and stem of Jacaranda 261 decurrens [26], roots of Chlorophytum borivilianum [27] and leaves of mangrove plants [28] 262 have been reported to produce hydrolytic enzymes of diverse types. All the aerobic 263 endophytic isolates of *H. spinosa* possessed catalase responsible for the decomposition of 264 hydrogen peroxide to less reactive oxygen and water molecules. Production of hydrolytic 265 enzymes gelatinase, amylase and lipase (Table 3) supports earlier observations [25-28]. The 266 presence of nitrate reductase and tryptophanase in some of the isolates appears to play a 267 key role in the nitrogen cycle and has important agricultural, environmental and public health 268 implications. The emergence of antibiotic resistance is not only limited to pathogenic 269 microorganisms but also found amongst environmental isolates as a result of horizontal 270 transfer of antibiotic resistance genes. Majority of the endophytes from H. spinosa showed 271 resistance to amoxycillin and bacitracin (Table 5) similar to those encountered in bacterial 272 endophytes of *P. foetida* [17], *Andrographis paniculata* [29] and mangrove plants [28].

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274 In view of the ever increasing demand for novel antimicrobial substances, the endophytes 275 have been identified as a potential source of antibiotics [6]. Several reports on the 276 antimicrobial evaluation of endophytic fungi from medicinal plants have been presented [30-277 32], antimicrobial activities of endophytic bacteria are not uncommon [17, 20, 29]. Li et al. 278 [30], however, have explored endophytic actinomycetes associated with pharmaceutical 279 plants in rainforest of Yunnan, China and detected endophytic Streptomyces displaying antimicrobial activities against S. aureus, E. coli and C. albicans. In the present study, nine 280 281 bacterial endophytes out of 11 from H. spinosa showed antibacterial activity against B. subtilis, B. cereus, E. coli, P. cepacia, K. pneumoniae and S. aureus following cross-streak assay (Table 6) and two of them showed broad spectrum antimicrobial activity indicating possible biotechnological potential. However, isolation, purification and detection of active compounds is in progress for their further utilization.

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287 **4. CONCLUSION**

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Endophytic bacterial isolates was found to be associated with leaves, stem and root of the medicinal plant, *H. spinosa* and they differed significantly in their morphological, physiological and biochemical characters. The endophytes also produced several hydrolytic enzymes of commercial importance. Antimicrobial evaluation of these culturable endophytes of *H. spinosa* has shown that they possess antibacterial activity against various bacterial species. The endophytes of traditional medicinal plants appear to be a potential source of antimicrobial metabolites as well as enzymes.

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302 COMPETING INTERESTS

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304 Authors have declared that no competing interest exists in performing this research and 305 preparation or publication of the results.

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308 AUTHORS' CONTRIBUTIONS

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This research was carried out through joint collaboration of authors AP and AKP. The authors took equal responsibilities to design the study, writing protocols, literature survey, performing the experiments and preparation of manuscript.

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