

A search for antiplasmodial metabolites among fungal endophytes of terrestrial and marine plants of southern India

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Abstract

Eighty four different fungal endophytes isolated from sea grasses (5), marine algae (36) and leaves or barks of forest trees (43) were grown *in vitro* and the secondary metabolites secreted by them were harvested by immobilizing them on XAD beads. These metabolites were eluted with methanol and screened using SYBR Green I assay for their antiplasmodial activity against blood stage *Plasmodium falciparum* in human red blood cell culture. Our results revealed that fungal endophytes belonging to diverse genera elaborate antiplasmodial metabolites. A *Fusarium* sp. (580, IC₅₀: 1.94 μ g ml⁻¹) endophytic in a marine alga and a *Nigrospora* sp. (151, IC₅₀: 2.88 μ g ml⁻¹) endophytic in a tree species were subjected to antiplasmodial activity-guided reversed phase high performance liquid chromatography separation. Purification led to potentiation as reflected in IC₅₀ values of 0.12 μ g ml⁻¹ and 0.15 μ g ml⁻¹ for two of the fractions obtained from 580. Our study adds further credence to the notion that fungal endophytes are a potential storehouse for a variety of novel secondary metabolites vested with different bioactivities including some that can stall the growth of the malaria parasite.

Keywords

Endophytes, antiplasmodial metabolites, Plasmodium falciparum, antimalarial compounds, marine-derived fungi

Introduction

Malaria poses a major health threat in developing countries and annually more than 200 million new malaria cases are reported (WHO 2011), emphasising the urgent need for new drugs against malaria. In India, its management is challenging due to the country's huge population and large geographical area further compounded by rapid urbanisation (Shah et al. 2011) as well as the resistance developed by the malarial parasite to most of the existing classes of antimalarial drugs (Anderson 2009). Although artemisinin, a sesquiterpene lactone of the Chinese herb Artemisia annua L., and its synthetic analogues are very effective in controlling drug-resistant malaria (White 2008), a recent report indicates the emergence of artemisinin-resistant malarial parasite strains in western Cambodia and western Thailand (Cheeseman et al. 2012). Such wide spectrum drug resistance and the possibility of the existence of regional subpopulations of malarial parasites differing in sensitivity to different classes of drugs necessitates continuing the search for novel antiplasmodial pharmacophores for short- and long-term effective control of malaria (Anderson 2009). Natural products with antiplasmodial activity appear to be superior to purely synthetic ones because (a) natural metabolites have better prospects of becoming drugs (Ortholand and Ganesan 2004) and (b) the genes coding for these metabolites can be cloned and overexpressed to obtain appreciable quantities of chemically complex metabolites like artemisinin at affordable costs (Westfall *et al.* 2012; Paddon *et al.* 2013). In this context, endophytes, an ecological group of fungi may be good sources of antiplasmodial metabolites.

Endophytes are mostly ascomycetous fungi which reside inside living tissues of all major groups of plants without producing any visible disease symptoms. Endophytes of both land (Suryanarayanan *et al.* 2009; Weber 2009) and marine (Suryanarayanan *et al.* 2010; Aly *et al.* 2011) plants produce a vast array of secondary metabolites exhibiting diverse bioactivities including antimicrobial, antifungal, free radical scavenging, cytotoxic and anti-insect activities. Based on *ex vivo* inhibition of growth of *Plasmodium falciparum* in human red blood cell cultures, we report antiplasmodial activity (IC₅₀: <10 mg/ml) in 19 of the 84 endophyte isolates screened by us. Although some antimalarial compounds have been reported

to be produced by a few free living marine fungi (Chinworrungsee *et al.* 2001; Wright and Lang-Unnasch 2005), here we report that marine endophytes are also a potential source of antiplasmodial compounds.

Materials and Methods

Endophyte isolates used

We have used for our investigations of endophytes, five isolates recovered from seagrasses, 36 isolates from marine algae (both from the coast of Rameswaram, Tamil Nadu) and 43 isolates from leaves or barks of trees growing in the forests of the Western Ghats. All these isolates were maintained in Vivekananda Institute of Tropical Mycology (VINSTROM), Ramakrishna Mission Vidyapith, Chennai. The endophytes were isolated by surface sterilising the plant tissues with ethanol and sodium hypochlorite (Suryanarayanan et al. 2010, 2011; Murali et al. 2013). The following isolates have been deposited at the National Fungal Culture Collection of India (NFCCI), Agharkar Research Institute, Pune [Sordaria sp. 1 (65) – Accession Number (AC) 10342, Curvularia lunata (93) -(AC) 10334, Pestalotiopsis (455) - (AC) 10336, Curvularia tuberculata (516) - (AC) 2434] and Institute of Microbial Technology (IMTECH), Chandigarh, India [Nigrospora oryzae (151) – (AC) 8660, Pithomyces maydicus (152) – (AC) 8657, Chaetomium sp. 1 (154) – (AC) 8658, Aspergillus terreus (543) - (AC) 10346, Phomopsis sp. (562) - (AC) 10345, *Curvularia* sp. (566) – (AC) 10337, *Trichoderma* sp. (568) – (AC) 10344, Drechslera papendorphii (589) – (AC) 10338, Colletotrichum sp. (599) - (AC) 1033 and Curvularia tuberculata (606) – (AC) 10340].

Extraction of secondary metabolites

Each fungus was grown as a static culture in 60 ml potato dextrose broth contained in a 250 ml flask along with XADTM 16 amberliteTM resin (Rohm and Hass, Philadelphia, PA, USA) (400 mg /flask) at 26°C for 21 days. XAD-16 amberlite is a polystyrene resin that adsorbs small molecular weight metabolites produced by fungi and facilitates their easy extraction (Suryanarayanan et al. 2010). Three replicates were maintained for each fungus. After incubation, the mycelium was filtered and the XAD from all the three flasks was combined, washed in distilled water and air dried. These dried XAD beads were shaken with 12 ml methanol (in a 15 ml Falcon tube) for 45 min at 25°C. Aliphatic alcohols like methanol exhibit unique abilities to extract diverse molecules of widely different polarities. Thus methanol can extract polar, amphipathic and also fairly non polar molecules. Methanol's economics and relatively low boiling point also encourages the use of this solvent for the first round of extraction whose aim is to capture molecules of diverse polarities. Supernatants were collected by decantation. This process was repeated thrice resulting in a total volume of 36 ml per sample. Methanol was evaporated at 40°C in a Labconco RapidVap parallel evaporation system. The residue was dissolved in 0.2 to 0.4 ml of methanol, transferred to pre-weighed microfuge tubes and subjected to evaporation to dryness. The weight of each elute was determined (Table I) before making stock solutions (25 mg/ml) in dimethyl sulfoxide (DMSO).

In vitro cultivation of Plasmodium falciparum

Chloroquine-sensitive 3D7 strain of P. falciparum (Pf3D7) was maintained in continuous culture following the method of Trager and Jensen (1976) with minor modifications. Cultures were maintained in fresh O^{+ve} human erythrocytes suspended at 4% (v/v) haematocrit in complete medium {16.2 g/L RPMI 1640 (Sigma) containing 25 mM HEPES, 11.11 mM glucose, 0.2% sodium bicarbonate (Sigma), 0.5% Albumax I (Gibco), 45 µg/litre hypoxanthine (Sigma) and 50 μ g/litre gentamicin (Gibco)} and incubated at 37°C in a gas mixture consisting of 5% O_2 , 5% CO_2 , and 90% N_2 . The spent medium was replaced with fresh complete medium every day to propagate the culture. Giemsa-stained blood smears were examined microscopically to monitor parasitemia. Synchronized ring stage parasite was obtained by 5% sorbitol (w/v) treatment (Lambros and Vanderberg 1979).

Drug/Crude extract dilutions

Stock solutions of XAD extracts were prepared in DMSO and that of Chloroquine phosphate (CQ) (Sigma) was prepared with Milli-Q grade water. The required drug concentrations were achieved by diluting the stocks with culture medium. The solutions of drugs and extracts were placed in 96-well flat bottom tissue culture grade plates (Corning).

Assay for anti-plasmodial activity

For drug screening, SYBR green I based fluorescence assay was used (Smilkstein et al. 2004, Bennet et al. 2004). The ability of SYBR green to give strong fluorescence only in the presence of DNA forms the basis for its use to assess cell proliferation. The absence of nucleus in human red blood cells where the malarial parasite proliferates allows the use of SYBR green for the specific monitoring of the growth of malarial parasite. However, caution is required since this fluorescence-based method is prone to quenching effects from the test samples which yield (antiplasmodial) false positives. In order to be not misled by such effects, each of the extracts which showed promise in the SYBR green assay was validated using Giemsa stained microscopy based analysis. Conversely, fluorescent molecules in test extracts can in principle lead to enhanced fluorescence even as these extracts may be antiplasmodial in action. However such cases of false negatives

Fungus	Extract ID	Strain isolated from #	Recovery (mg/400 mg XAD)	<i>Pf3D7</i> IC ₅₀ μg ml ⁻¹	HeLa TC ₅₀ µg ml ⁻¹	Selectivity Index IC ₅₀ /TC ₅₀
Uninoculated growth medium	Control		18	>100		30 30
Alternaria sp.	572	Sargassum wightii (A*)	16.53	5	44	8.8
Alternaria sp.	590	<i>Turbinaria</i> sp. (A*)	16.52	5.2	67	12.9
Alternaria sp.	592	Turbinaria sp. (A)	31.47	16		
Aspergillus terreus	526	Caulerpa sertularioides (A)	40.54	43		
Aspergillus terreus	543	Ulva lactuca (A)	46.01	37		
Aspergillus niger	544	Ulva lactuca (A*)	17.64	1.2	25	20
Aspergillus sp.	551	Halimeda macroloba (A)	14.59	18		
Aspergillus sp. 2	650	Padina tetrastromatica (A)	34.51	100		
Aspergillus janus	581	Sargassum wightii (A)	14.49	24		
Aspergillus sp. 4	597	<i>Turbinaria</i> sp. (A)	26.41	26		
Chaetomium sp.	542	Gracilaria edulis (A*)	11.25	2.6	35	13.5
Chaetomium sp.	548	Ulva lactuca (A*)	19.78	4.8	77	16
Chaetomium sp.	558	Padina tetrastromatica (A)	6.91	85		
Chaetomium sp.	628	Grateloupia lithophila (A)	3.65	15		
Cladosporium sp.	520	Caulerpa racemosa (A)	7.97	>100		
Cladosporium sp.	564	Portieria hornemannii (A)	7.98	21		
<i>Curvularia</i> sp.	566	Sargassum wightii (A)	31.01	40		
Curvularia lunata	602	Sargassum sp. (A)	12.76	14		
Curvularia tuberculata	606	<i>Turbinaria</i> sp. (A)	18.81	44		
Colletotrichum sp.	599	Sargassum sp. (A)	4.66	45		
Drechslera sp.	589	<i>Turbinaria</i> sp. (A)	11.46	39		
Drechslera sp.	610	Turbinaria sp. (A)	6.71	40		
Emericella nidulans	560	Halymenia sp. (A)	9.36	36		
Fusarium sp.	580	Sargassum wightii (A*)	42	2	72	36
Memnoniella sp.	648	Portieria hornemannii (A)	26.39	60		
Nigrospora sp.	546	Ulva lactuca (A)	15.65	17		
Nigrospora sp.	594	<i>Turbinaria</i> sp. (A)	2.9	13		
Paecilomyces sp.	534	Caulerpa scalpelliformis (A*)	14.73	9.7	70	7.2
Penicillium sp.	536	Caulerpa scalpelliformis (A)	51	43		
Penicillium sp.	552	Halimeda macroloba (A*)	8.43	4.8	>100	>20
Pestalotiopsis sp.	574	Sargassum wightii (A)	7.54	31		
Phoma sp.	605	Sargassum sp. (A)	8.56	25		
Phomopsis sp.	562	Portieria hornemannii (A)	12.58	39		
Pithomyces sp.	604	Sargassum sp. (A)	11.74	46		
Trichoderma sp.	568	Sargassum wightii (A*)	25.13	1.56	100	64.1
UNH-1	649	Padina sp. (A)	23.73	34		
Alternaria sp.	105	Bridelia retusa (E)	25.97	21		
Arthrinium sp.	87	Elaeocarpus serratus (E)	15.34	>100		
Auerobasidium pullulans	223	Spinifex littoreus (E)	14.74	71		
Botrytis sp.	69	Erythroxylon monogynum (E)	38.74	100		
Chaetomium sp.	154	<i>Phoebe lanceolata</i> (E)	18.38	22.5		
Chaetomium sp.	672	Phoebe lanceolata (E*)	19.4	3.5	80	22.9
Cladosporium sp.	670	Bridelia retusa (E)	5.74	>100		
Cladosporium sp.	675	Elaeocarpus serratus (E)	8.17	>100		
Curvularia lunata	93	Ilex wightiana (E)	5.43	>100		
Curvularia sp.	100	Macaranga peltata (E)	24.21	31		

Table I. Antiplasmodial potencies, mammalian cell cytotoxicities and selectivity indices of endophytic fungal secretions

Fungus	Extract ID	Strain isolated from #	Recovery (mg/400 mg XAD)	<i>Pf3D7</i> IC ₅₀ μg ml ⁻¹	HeLa TC ₅₀ μg ml ⁻¹	Selectivity Index IC ₅₀ /TC ₅₀
Curvularia sp.	104	Rhizophora mucronata (E)	4.42	12.5		
Curvularia sp.	456	Acacia leucophloea (E*)	27.21	2.5	50	20
Curvularia eragrostidis	669	<i>Litsea floribunda</i> (E)	14.25	78		
Colletotrichum sp.	85	Catharanthus roseus (E)	12.02	97		
Colletotrichum gloeosporioides	90	Cinnamomum wightii (E)	14.84	14		
Corynespora sp.	668	<i>Kydia calycina</i> (E*)	8.11	5	>100	>20
Corynespora sp.	673	<i>Murraya paniculata</i> (E)	4.76	>100		
Drechslera sp.	83	Catharanthus roseus (E)	18.64	20		
Fusarium sp.	78	Azadirachta indica (E)	13.6	75		
Fusarium sp.	204	Euphorbia tithymaloides (E)	5.02	28		
Fusarium sp.	217	Vitex altissima (E*)	3.79	1.56	30	
Fusarium sp.	467	Semecarpus anacardium (E)	3.97	45		
Fusarium sp.	674	<i>Murraya paniculata</i> (E)	4.91	>100		
Lasiodiplodia theobromae	454	Bridelia retusa (E)	7.49	49		
Lasiodiplodia theobromae	671	Randia dumetorum (E)	9.37	80		
Nigrospora oryzae	76	Cryptocarya bourdillonii (E)	3.68	>100		
Nigrospora oryzae	97	Daphniphyllum neilgherrense (E)	6.09	>100		
Nigrospora oryzae	151	Macaranga peltata (E*)	21.97	3	>100	>33
Nodulisporium sp.	443	Syzygium densiflorum (E)	5.52	77		
Nodulisporium sp.	95	Glochidion sp. (E)	12.03	75		
Penicillium sp.	141	<i>Phyllanthus emblica</i> (E)	12.44	>100		
Pestalotiopsis sp.	117	<i>Litsea stocksii</i> (E)	4.75	>100		
Pestalotiopsis sp.	455	Bridelia retusa (E)	16.06	75		
Phomopsis sp.	676	<i>Murraya paniculata</i> (E)	12.78	11	52	4.8
Phomopsis sp.	677	Lagestroemia microcarpa (E*)	10.67	8	100	12.5
Pithomyces sp.	152	<i>Eurya nitida</i> (E)	11.3	>100		
Phyllosticta capitalensis	666	Maytenus emarginata (E)	7.82	>100		
<i>Sordaria</i> sp.	65	<i>Eurya nitida</i> (E)	13.86	>100		
<i>Xylaria</i> sp.	123	Anogeissus latifolia (E)	12.32	>100		
<i>Xylaria</i> sp.	184	Randia dumetorum (E)	13.4	>100		
<i>Xylaria</i> sp.	195	Ziziphus jujuba (E)	44.26	81		
<i>Xylaria</i> sp.	315	Butea monosperma (E)	8.85	>100		
<i>Xylaria</i> sp.	667	Lasianthus venulosus (E)	23.74	3.8	<50	<13
Aphanocladium sp.	508	Thalassia sp. (Sg)	9.41	97		
Auerobasidium pullulans	585	Thalassia sp. (Sg)	20.16	93		
Curvularia tuberculata	516	Syringodium sp. (Sg)	15	33		
Gonatophragmium mori	622	Thalassia sp. (Sg)	8.13	37		
Memnoniella sp.	647	Cymodocea serrulata (Sg)	30.45	67		

[#] A = marine algal endophytes, E = tree endophytes, Sg = seagrass endophytes, * indicates strains selected for sequencing of fungal ITS region

were not observed in the extracts studied by us. Sorbitol-synchronized ring stage parasites (haematocrit: 2%, parasitaemia: 1%, 100 μ l) under normal culture conditions were incubated in the presence or absence of increasing concentrations of the extracts. It is important to note that use of synchronized cultures over mixed stage cultures is necessary for (a) consistency in results across laboratories, (b) use of ring stage synchronized cultures can enable the test molecules to interact with all the three stages (ring, trophozoite and schizont) of the 48 hrs long life cycle of *P. falciparum* in culture, (c) starting the experiment with synchronized ring stage culture provides the distinct advantage of observing growth inhibitory effects without a rise in parasitemia during the ring-trophozoite-schizont transitions and (d) the choice of ring stage synchronized cultures is also governed by the procedural ease (5% Sorbitol treatment which destabilizes both trophozoites and schizonts

without affecting ring stage cells) with which pure rings can be obtained against the elaborate Percoll gradient based centrifugal procedures required to obtain either the trophozoites or the schizonts.

CQ was used as positive control and 0.4% DMSO (v/v), which was found to be non-toxic to the parasite, was used as vehicle control. After 48 h of incubation, 100 µl of SYBR Green I buffer [0.2 µl of 10,000 × SYBR Green I (Invitrogen) per ml of lysis buffer {Tris (20 mM; pH 7.5), EDTA (5 mM), saponin (0.008%; wt/vol), and Triton X-100 (0.08%; vol/vol)}] was added to each well, mixed twice gently with multi-channel pipette and incubated in the dark at 37°C for 1 h. Fluorescence was measured using a Victor fluorescence multi-well plate reader (Perkin Elmer) with excitation and emission at 485 and 530 nm, respectively. Fluorescence counts for CO (0.1 μ M for *Pf*3D7) were deducted from counts in each well. A dose-response curve was constructed by plotting fluorescence counts against the drug concentration and IC_{50} (dose of a drug required to slow the rate of growth of a cell population by 50%) was determined. In the experiments described by us, no drug control corresponds to 100 % growth while 100 nM Chloroquine (sufficient to cause total arrest of growth) corresponds to 0 % growth. Between these two boundaries of 100% and 0% growth lie the points where at different concentrations of test samples, the parasite shows different extents of growth. The graph depicting growth inhibition as a function of concentration of test sample is used to obtain IC_{50} . Giemsa-stained smears of extract-treated parasite cultures were visualized microscopically to validate the results from fluorescence-based assay.

Cytotoxic activity of extracts on HeLa cells using MTT assay

The cytotoxic effects of active extracts on mammalian cells was determined by a functional assay (Mosmann 1983) using HeLa/L929 cells cultured in complete medium containing 16.2 g/l RPMI 1640, 10% fetal bovine serum, 0.21% sodium bicarbonate (w/v) (Sigma) and 50 µg/mL gentamycin. Cells (10⁴ cells/200 µl/well) were seeded into 96-well flat-bottom tissue culture plates in complete medium. After 24 h of seeding, test solutions were added and cells incubated for 48 h in a humidified atmosphere at 37°C and 5% CO₂. DMSO (as positive inhibitor) was added at 10% v/v. Twenty microlitres of a stock solution of MTT [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide] (5 mg/mL in 1X phosphate buffered saline) was added to each well, gently mixed and incubated for another 4 hr. After centrifugation at 300 g for 5 min, the supernatant was removed and 100 µl DMSO (quench agent) was added to the cell pellet. Formazan formation was read using a microtiter plate reader (VersaMax tunable multiwell plate reader) at 570 nm. The 50% cytotoxic concentration (TC₅₀) of drug was determined by analysis of dose-response curves. Selectivity index (TC_{50}/IC_{50}) was also calculated for the different extracts studied.

Bioassay Guided Fractionation of Potent Extracts

Following the screening of 84 different fungal XAD extracts, the extracts of isolate numbers 151 and 580 which showed promising antiplasmodial potency ($IC_{50} \leq 3 \mu g/ml$) and selectivity indices (>30) were chosen for antiplasmodial activity guided reversed- phase HPLC fractionation. Waters C18, Deltapak (19 × 300 mm, 15 μ) column was used for fractionation using methanol water gradient (1%/min) at a flow of 10 ml/min on Gilson prep HPLC system. Dual wavelength detection at 214 nm (σ - σ * transitions shown by several aliphatic molecules) and 254 nm (π - π * transition shown by aromatic molecules) is helpful in assessing the chemical nature of molecules eluting from the column. Fractions were collected, dried, weighed and analyzed for antiplasmodial and cytotoxic activity.

Fungal DNA extraction

The taxonomic affiliation of seven fungal isolates which showed relatively high antiplasmodial activity was confirmed by sequencing of ITS1-5.8S-ITS2 region. Genomic DNA was extracted from pure cultures of fungal endophytes following the phenol-chloroform extraction method. For this, cultures were grown on potato dextrose agar plates for 7 days and mycelia were collected and transferred to a sterile microfuge tube containing 500 µL of DNA extraction buffer (0.1 M NaCl, 50 mM Tris, 10 mM Na, EDTA, 2% SDS, pH 8.0). The samples were then ground well and to this emulsion, 500 μ L of chilled phenol was added and centrifuged at 15,300g for 15 min at 4°C. The upper aqueous phase was transferred to a fresh tube and an equal volume of chilled chloroform: isoamylalcohol (24:1 v/v) was mixed to get a suspension. The tubes were centrifuged at 15,300g for 15 min-at 4°C and the clear aqueous phase was transferred to a fresh microfuge tube. After adding chilled isopropanol (0.6 volume) and 3 M sodium acetate (0.1 volume), the contents were gently mixed and incubated at -80°C for 2 h followed by centrifugation at 15,300g for 15 min at 4°C. The supernatant was discarded and the pellet was washed with 200 μ L of 70% ethanol (v/v) followed by centrifugation at 15,300g for 10 min at 4°C. After discarding the supernatant, the pellet was air-dried, dissolved in 50 μ L of sterile distilled water and stored at -20°C. The genomic DNA was electrophoresed in 0.8% agarose gel (w/v) and then diluted suitably for amplifying the nuclear ITS region.

PCR amplification and sequencing of fungal ITS region

The nuclear ribosomal internal transcribed spacers (ITS) along with 5.8S region were PCR amplified using primer sets ITS5 and ITS4 or ITS1F and ITS4 (White *et al.* 1990; Gardes and Bruns 1993). A 25 μ L amplification reaction consisted of 2.5 μ L dNTPs (10 μ M), 2.5 μ L PCR buffer (10X), 1 μ L primers (forward and reverse – 10 μ M each), 1 μ L Taq DNA Polymerase (1 U), 1 μ L DMSO (1%, v/v), 1 μ L MgCl, (25 mM) and 1 μ L fun-

gal genomic DNA as template (approx. 25–50 ng). Cycling reactions were performed in a Master Cycler Thermocycler (Eppendorf, USA) with following conditions: 94 °C for 3 min, followed by 34 cycles of 94 °C for 30 s, 54 °C for 30 s, 72 °C for 1 min; and finally 72 °C for 10 min. The amplified products were run on 1% agarose gel (w/v) to determine the size and purity of amplicons. The purified PCR products were then sequenced in an automated sequencer (ABI 3130 Genetic Analyzer). The sequences obtained were manually edited before performing a BLAST search using the megablast algorithm to arrive at the nearest match. The results were used to confirm the identity of fungal morphotypes at species level wherever possible. The nucleotide sequence data obtained were deposited in GenBank database (Accession numbers KF135618-KF135624).

Phylogenetic analysis

The nuclear ITS1-5.8S-ITS2 sequences obtained for different endophytes isolated in the present study were aligned with 250 related sequences available in the GenBank database based on the results obtained from BLAST search. The sequences were then aligned by ClustalW (Thompson *et al.* 1994) using the default settings and manually re-edited to build an initial neighborjoining analysis. From this dataset, we selected 89 closely related sequences to perform the final analysis. The neighbour joining method of Saitou and Nei (1987) was used to construct the evolutionary tree from the datasets by using MEGA version 5.1 (Tamura *et al.* 2011). The evolutionary distance was calculated using the Maximum Composite Likelihood method and branch support was calculated with average distances from 1,000 bootstrap replicates (Felsenstein 1985). All gaps and missing data in the dataset were eliminated for this analysis.

Results

Antiplasmodial potencies, mammalian cell cytotoxicities and selectivity indices of endophytic fungal secretions

Sixteen endophyte isolates belonging to twelve different fungal genera (Alternaria, Aspergillus, Chaetomium, Curvularia, Corynespora, Fusarium, Nigrospora, Paecilomyces, Penicillium, Phomopsis, Trichoderma and Xylaria) which were isolated from marine algae and angiosperm trees, representing about 19% of the 84 fungi screened, showed high (IC_{50} <10 μ g/ml) antiplasmodial activity (Table I). The antiplasmodial IC₅₀ potency profile of the remaining XAD elutes was 11–25 µg/ml (18% isolates), 26-50 µg/ml (21% isolates), 51-100 μ g/ml (19% isolates) (Fig. 1). The growth curves obtained using the SYBR Green assay for two representative XAD extracts viz. 151 (IC₅₀ 2.88 μ g/ml) and 580 (IC₅₀ 1.94 μ g/ml) together with Giemsa stained smears for control and treated samples are shown in Fig. 2. The correspondence between % growth as scored in SYBR Green assay and % parasitemia in microscopic images provides validity to the results obtained from the SYBR Green assay. Furthermore, the extracts which showed $IC_{50} \le 11$ µg/ml were analysed for their selectivity against Plasmodium over mammalian cells. Most of these extracts were found to be nontoxic with selectivity index of >10 (Table I).

Antiplasmodial activity guided RPHPLC fractionation of extracts 151 and 580

The extracts of *Fusarium* sp. (580) endophytic in a marine alga and *Nigrospora* sp. (151) endophytic in a tree species which showed the best combination of the features of high yield



Fig. 1. Frequency of fungal endophytes displaying different antiplasmodial potencies as IC_{s0} in μ g ml⁻¹ obtained after screening eighty four different fungal secretions captured on XAD resin. The relative percentage of endophyte strains within each potency range is indicated in parentheses



Fig. 2. *P. falciparum* 3D7 was *ex vivo* cultured for 48 hrs in human red blood cells in absence and presence of increasing concentrations of XAD extracts of 151 and 580 respectively. Graphs show % growth measured by SYBR Green fluorescence assay as a function of increasing concentrations of extracts. Each data point represents the mean +/- standard deviation of three independent observations. ICEstimator-version 1.2 (http://www.antimalarial-icestimator.net/MethodIntro.htm) was used for plotting the graph and obtaining the IC_{50} values. Microscopic images of Giemsa stained smears of untreated vs XAD extract (taken at IC_{100}) treated cultures are shown. The high ring stage parasitemia in control vs the low moribund trophozoite stage parasitemia in extract treated cultures provides validity to the SYBR Green results

(weight of metabolites obtained on elution from XAD beads), antiplasmodial potency and selectivity index were subjected to RPHPLC based activity guided fractionation. As shown in Fig. 3 and Fig. 4, the fractionation of crude extracts led to identification of fractions with high activity. A marked increase in potencies of crude (IC_{50} : 2 µg ml⁻¹) vs fractions 6 & 7 (IC_{50} : 0.12 µg ml⁻¹ and 0.15 µg ml⁻¹) was observed in the case of 580 (Fig. 4).

Phylogenetic study

The seven representative fungal isolates which showed low IC_{50} values (IC_{50} <4.8 µg ml⁻¹) were selected and studied using molecular techniques to infer their phylogeny. The

ITS region was amplified and sequenced using fungal specific primers and the sequences obtained were compared with GenBank (using megablast algorithm) and CBS databases to identify the closest matching sequences (Table II). The search results showed that the seven isolates belonged to six different genera namely, *Alternaria, Chaetomium, Curvularia, Fusarium, Nigrospora* and *Trichoderma*. However, we found that the top match with reference to species names from two different databases varied for three strains (strains 456, 151, 590). Isolate 580 (marine algal endophyte) showed 100% match with *Fusarium oxysporum* f. sp. *aechmeae* and isolate **151** (tree endophyte) showed a 100% match with *Nigrospora sphaerica* (Table II). The discrep-



Fig. 3. Antiplasmodial activity guided Semiprep Reversed Phase HPLC Chromatogram of *Fusarium* sp. (580) XAD extract. Supernatant (32 mg/ml DMSO) was injected to Deltapak (C18, 19×300 mm, 15μ) column and fractionated using Methanol – water (10 ml/min, 1%/min) gradient (slanting line). Absorbance at 254 nm (lower chromatogram) and 214 nm (upper chromatogram) are shown. Fractions (Frc, 1–11 indicated by filled and empty bars) were collected, dried, weighed and analyzed for antiplasmodial and HeLa cells cytotoxic activity. Inset shows amount (mg) recovered, antiplasmodial IC₅₀, cytotoxic activity and selectivity index (SI). Note the significant enhancement in antiplasmodial potency and selectivity indices of fractions 6 and 7 in comparison with the activities for the crude extract (Cr)

ancy in species level identification for the three strains can be attributed to the fact that more than 10% of sequences available in public sequence databases may have incorrect species annotation (Nilsson et al. 2006; Ko et al. 2011). In our study, we have carried out morphological characterization of these strains to further validate the results obtained from molecular characterization. Furthermore, we selected sequences related to these seven strains from the GenBank database (Table III) based on the results obtained from BLAST search to construct a neighbour joining tree. The sum of branch length of optimal tree constructed using the maximum composite likelihood approach was found to be 0.667. The phylogram showed that the endophyte isolates and the related sequences were grouped into six (A-F) major clades (Fig. 5) with all clades receiving very high branch support. Clades (A-F) represented the isolates belonging to species of Fusarium, Nigrospora, Trichoderma, Chaetomium, Alternaria, and Curvularia respectively. Interestingly, however, a few of the Alternaria species grouped together with species belonging to Curvularia. Two of the isolates (542 and 548) could be grouped together with Chaetomium species in clade D.

Discussion

Many recent studies have shown that endophytes produce an extraordinary array of functional metabolites (Schueffler and Anke 2011; Tejesvi and Pirttila 2011). This need for such a diversity of metabolites may be triggered by the fact that endophytes which survive in living plant tissues may need them for their constant interaction with the defence chemicals produced by the plant host (Schulz et al. 1999). It is quite likely that survival in an alien life form may have resulted in the evolution of heightened synthetic ability of endophytes (Suryanarayanan et al. 2009). The hypothesis that horizontal gene transfer events from the host plant to its endophytes being instrumental in the production of some of these metabolites has evoked considerable interest as well as great controversy (Heinig et al. 2013; Sachin et al. 2013). Pharmaceutically important secondary metabolites-producing endophytes, compatible with in vitro cultures offer special advantages that include cost- effective large scale production of the metabolites in industrial laboratory settings. In this regard, endophytes from marine habitats appear to be unique since many of them produce different novel



Fig. 4. Antiplasmodial activity guided Semiprep Reversed Phase HPLC Chromatogram of *Nigrospora* sp. (151) XAD extract. Supernatant (500 mg/2 ml methanol) was injected to Deltapak (C18, 19×300 mm, 15μ) column and fractionated using Methanol – water (10 ml/min, 1%/min) gradient (slanting line). Absorbance at 254 nm (lower chromatogram) and 214 nm (upper chromatogram) are shown. Fractions (Frc, 1–14 indicated by filled and empty bars) were collected, dried, weighed and analyzed for antiplasmodial and HeLa cells cytotoxic activity. Inset shows amount (mg) recovered, antiplasmodial IC⁵⁰, cytotoxic activity (TC₅₀) and selectivity index (SI), crude extract (Cr)

Table II. Fungal cultures with their closest matches from GenBank and CBS	databases based on ITS sequence similarity. Numbers in paren-
thesis indicate the accession number of the closest matched sequences	

Culture Code	Identified as	Top match in GenBank	Similarity	Top match in CBS	Similarity
542	Chaetomium sp.	Chaetomium spirochaete (JN209921)	100%	Chaetomium spirochaete (CBS730.84)	100%
456	Curvularia sp.	Cochliobolus nisikadoi (JN943428)	98%	Curvularia ischaemi (CBS630.82)	97%
151	Nigrospora sp.	<i>Nigrospora</i> sp. (HQ631070)	99%	Nigrospora sphaerica (CBS167.26)	100%
590	Alternaria sp.	Alternaria brassicae (JX984695)	99%	Alternaria rhizophorae (CBS118816)	99%
580	Fusarium sp.	Fusarium oxysporum (KC254033)	100%	<i>Fusarium oxysporum</i> f. sp. <i>aechmeae</i> (CBS244.61)	100%
568	Trichoderma sp.	Trichoderma harzianum (KC330218)	100%	Trichoderma harzianum (CBS354.33)	100%
548	Chaetomium sp.	Chaetomium globosum (JF826006)	100%	Chaetomium globosum (CBS 105.40)	100%

Table III. Sequences of fungal species used for constructing the phylogram

Species	GenBank Accession Number	Species	GenBank Accession Number	
Alternaria aff. longipes	DQ156337	Cochliobolus australiensis	JN943408	
Alternaria alternata	GQ916545	Cochliobolus cymbopogonis	JQ783057	
Alternaria arborescens	JQ676197	Cochliobolus geniculatus	AB245085	
Alternaria brassicae	JX857165	Cochliobolus lunatus	HQ607991	
Alternaria californica	JQ693645	Cochliobolus nisikadoi	JN943428	
Alternaria dianthicola	JQ693640	Curvularia affinis	GU073105	
Alternaria frumenti	JQ693654	Curvularia clavata	GQ179976	
Alternaria hordeicola	JQ693642	Curvularia coicicola	AB453880	
Alternaria humuli	JQ693652	Curvularia fallax	JQ360963	
Alternaria incomplexa	JQ693658	Curvularia geniculata	GU073454	
Alternaria intercepta	JQ693656	Curvularia inaequalis	HM101095	
Alternaria merytae	JQ693651	Curvularia intermedia	GU073103	
Alternaria metachromatica	JQ693660	Curvularia pseudorobusta	AB453879	
Alternaria novae-zelandiae	JQ693655	Curvularia sichuanensis	AB453881	
Alternaria panax	JQ693662	Curvularia trifolii	GQ241277	
Alternaria porri	HM204456	Fusarium cf. solani	JX270188	
Alternaria rosae	JQ693639	Fusarium equiseti	HQ649908	
Alternaria solani	JX469421	Fusarium incarnatum	JX885463	
Alternaria sp.	KC010550	Fusarium oxysporum	KC254033	
Alternaria tenuissima	JX860514	Fusarium oxysporum	JX885462	
Chaetomium arcuatum	AB746177	Fusarium oxysporum f. sp. cyclaminis	JQ676177	
Chaetomium atrobrunneum	JN034195	Fusarium oxysporum f. sp. melonis	GU934523	
Chaetomium brasiliense	FR718872	Fusarium oxysporum f. sp. ranunculi	JQ340086	
Chaetomium cancroideum	HM449046	Fusarium pseudoanthophilum	HF548703	
Chaetomium carinthiacum	HF548694	Fusarium sp.	HQ130709	
Chaetomium coarctatum	HM365260	Gibberella moniliformis	JQ277275	
Chaetomium cruentum	HM365266	Hypocrea lixii	JX436467	
Chaetomium dolichotrichum	HM449049	Hypocrea nigricans	JN943372	
Chaetomium elatum	JN209874	Hypocrea rufa	AY380908	
Chaetomium erectum	HM449044	Hypocrea stilbohypoxyli	AY380916	
Chaetomium funicola	AB746176	Hypocrea virens	JX969615	
Chaetomium globosum	JF826006	Nigrospora cf. sphaerica	JQ676183	
Chaetomium gracile	HF548698	Nigrospora oryzae	EU821485	
Chaetomium grande	HM365253	Nigrospora sp.	HQ631070	
Chaetomium murorum	GQ376100	Nigrospora sp.	JF694932	
Chaetomium nigricolor	AB746178	Nigrospora sphaerica	GQ258792	
Chaetomium nigricolor	AJ458185	Trichoderma aureoviride	FJ487919	
Chaetomium ochraceum	JN093258	Trichoderma cf. harzianum	KC176363	
Chaetomium piluliferum	HE649377	Trichoderma harzianum	KC330218	
Chaetomium reflexum	HM449051	Trichoderma ovalisporum	AY380896	
Chaetomium sphaerale	AB625588	Trichoderma piluliferum	JQ517493	
Chaetomium spirochaete	JN209921	Trichoderma pleuroticola	HM142362	
Chaetomium subaffine	HM365247	Trichoderma pleurotum	HM142363	
Chaetomium undulatulum	HM365250	Trichoderma sp.	FJ645728	
		Trichoderma tomentosum	FJ487916	



Fig. 5. Neighbour joining tree for sequences obtained from the present study (marked as dark circles) along with 89 other related sequences obtained from GenBank database. The evolutionary distance was computed following the Maximum Composite Likelihood method. Bootstrap values above 75% are shown as numbers above nodes of branches. Sequences used in the tree construction are listed in Table III

metabolites (Bugni and Ireland 2004; Jones *et al.* 2008; Raghukumar 2008; Schulz *et al.* 2008; Kjer *et al.* 2010; Suryanarayanan *et al.* 2011; Flewelling *et al.* 2013). Nearly 30% of the novel metabolites produced by marine-derived fungi are from fungi associated with marine algae (Bugni and Ireland 2004); some of these fungi produce metabolites having hitherto unknown carbon frameworks (Kjer *et al.* 2010). This underscores the importance of our study with the secretions from *in vitro* grown endophytes of not only angiosperms but also seagrasses and seaweeds that exhibit potent anti *P. falciparum* activities.

Although only a limited number of endophyte isolates were screened in the present study, many of them produced antiplasmodial compounds; the finding that as high as 19% and 18% of the screened fungal isolates showed IC₅₀<10 μ g/ml and IC₅₀ of 11–25 μ g/ml, respectively, suggests that deeper exploration of endophytic fungi may reveal a rich repertoire of antiplasmodial molecules. Our findings have in-

dicated that endophytic fungi belonging to the genera Alternaria, Aspergillus, Chaetomium, Curvularia, Corynespora, Fusarium, Nigrospora, Paecilomyces, Penicillium, Phomopsis, Trichoderma and Xylaria should be studied intensely for novel antiplasmodial compounds. With reference to the identification of the fungal isolates, we could confirm only 4 of our seven endophyte isolates to the species level based on the similarity search from two different databases. This could be attributed to the fact that nearly 50% of the ITS sequences in the databases are without a species name (Benson et al. 2010) and nearly 10% of them have incorrect names (Nilsson et al. 2006). We used ITS sequences for characterising the fungi taxonomically as this region has been used widely to infer phylogeny and species identification in fungi (Begerow et al. 2010). ITS sequences have been found to be a better marker for fungi than even the genes for ribosomal large and small subunits, since this region has good species resolution capabilities (Schoch et al. 2012).

Although a few terrestrial plant endophytes have been shown to have antiplasmodial activity (Kongsaeree et al. 2003; Cao et al. 2011), our results show that endophytes colonizing seaweeds are equally a potential source to provide new leads against malaria. Further, our data suggest that antiplasmodial potential is more widespread across diverse fungal genera. Our attempts at antiplasmodial activity-guided RPHPLC purification of secretions from isolates 151 and 580 have led to the segregation of fractions with low and high activity. Indeed, in the case of 580 (IC₅₀ crude 1.94 μ g/ml), we were able to obtain fractions with IC_{50} down to 120 ng/ml. We are optimizing the production of such potent antiplasmodial metabolites from the identified endophytes to be followed by their isolation and identification for the exploration of their potential as novel drugs against malaria. Additionally, we are also examining the qualitative and quantitative changes in the antiplasmodial metabolites spectrum of isolates 151 and 580 as influenced by culture conditions since culture conditions are known to induce production of several new metabolites by fungi (Bode et al. 2002; OBrian et al. 2007).

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