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Antimycobacterial activity of cyanobacterial species isolated from the coastal regions of Tamil Nadu

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The extract of *Geitlerinema carotinosum* CNP 4003 showed promising antimicrobial activity (100 and 1000 μ g/disc/well) against *E. coli* ATCC 35218, *S. aureus* ATCC 25923, and *M. smegmatis* with inhibition zones of 6, 8, and 11 mm, respectively. In addition, the extract also exhibited cytotoxic activity (IC₅₀ = 175 μ g/ml) against mammalian (HepG2) liver cancer cell lines. Therefore, the crude extract was fractionated using column chromatography technique and the active fractions were identified. The active fractions tested against HepG2 cell line showed 95 % hemolytic activity at the concentration of 375 μ g/ml with IC₅₀ as 63 μ g/ml. Further, the compounds in the active fractions were analyzed and the results indicated the presence of indoles, terpenes, and peptides. The chemical composition of the active fraction was analyzed by using gas chromatography-mass spectrometry (GC-MS). The potential antimycobacterial strain G. *carotinosum* CNP 4003 was confirmed by molecular characterization and the DNA sequence was deposited in the gene bank.

[Keywords: Cytotoxicity, Geitlerinema sp., Hemolytic, Marine Cyanobacteria, Mycobacterium smegmatis]

Introduction

Tuberculosis (TB) is precarious and а communicable bacterial infection that causes considerable and public socioeconomic health problems in developing countries. Members of the Mycobacterium species such as Mycobacterium bovis, M. africanum, M. tuberculosis, M. pinnipedii, M. canettii, and M. microti are the main causative agents of TB causing significant health losses in human beings¹. Around the world, *M. tuberculosis* is the predominant species to cause TB. M. tuberculosis is frequently developing resistance against all first- and second-line drugs like isoniazid, rifampicin, ethambutol, streptomycin, pyrazinamide, aminoglycosides, and fluoroquinolones². Therefore, it is a vital requirement to find out novel medicines through antimycobacterial action valuable which is commercially viable, and eco-friendly. Various infectious diseases are known to be treated by natural products throughout the history of mankind³.

Among the mycobacterium species, *M. smegmatis* is classified as saprophytic, relatively safe, and incapable of causing disease and doesn't usually cause disease in any humans⁴. Further, due to its properties such as non virulence, fast grower or fast doubling time and

similarities with *M. tuberculosis* (> 2000 common homologs; unusual cell wall structure that can oxidize carbon monoxide aerobically), it has become the simple model for TB. In modern times, several new species belonging to this genus (*M. avium*, *M. leprea*, *M. bovis*, and *M. tuberculosis*) are identified as virulent that cause TB and leprosy. Therefore, non virulent bacteria of this type are necessary to tackle the mycobacerium associated health problems. In the majority cases, *M. smegmatis* is used as secure and a virulent alternative⁵. One of our earlier reports emphasized the efficacy of ethyl acetate extract (from the *Oscillatoria laetevirens* BDU 141071) activity against *M. smegmatis*⁶.

Extracts from *Spirulina platensis* isolated from an Egyptian water station has been accounted to hold anti-bacterial properties. Methanolic and aqueous extracts from the powder of cyanobacterial culture were found to possess action beside *Streptococcus faecalis*, *Salmonella typhimurium* and *Escherichia coli* 100 % (0.3 mg/ml), 86.2 % (0.5 mg/ml) and 91.6 % (0.7 mg/ml) from MeOH and 72.6 % (0.1 mg/ml), 99.3 % (0.9 mg/ml) and 74.4 % (0.9 mg/ml) from aqueous extracts)⁷. Murugan⁸ reported excellent activity of sodium phosphate buffer extracts of

S. platensis (C-phycocyanin) against Klebsiella pneumonia, Bacillus subtilis, E. coli, Staphylococcus aureus (11, 11, 6, and 12 mm of inhibition zone, respectively at 30 μ g/disc). Also, aqueous extracts of Oscillatoria sp. showed activity against Streptococcus mutants (20 mm of zone at 750 μ g/ml)⁹. Crude extracts of Lyngbya aestuarii and Oscillatoria boryana inhibited the growth of three of the pathogens, E. coli, Enterobacter aerogenes and Salmonella typhi¹⁰.

The hexane extract of Trichodesmium erythraeum exhibited antifungal activity against Trichophyton and simii (31.25) $\mu g/ml$) Τ. rubrum, T. mentagrophytes, Aspergillus flavus, A. niger, Scopulariopsis sp, and Botrytis cinerea at 1000 $\mu g/ml^{11}$. Ethyl acetate extract of *Phormidium* valderianum exhibited anticandidal activity at the concentration of 0.5 mg/disc against Candida albicans¹². Frankmölle et al.¹³ demonstrated a moderate antifungal action of crude ethanolic extracts of Anabaena laxa on Aspergillus oryzae and C. albicans (26 and 19 mm of inhibition zone at > 10Further, the methanolic $\mu g/ml$). extracts of L. aestuarii showed a reasonable antifungal action to C. $albicans^{14}$.

Freshwater, terrestrial and marine cyanobacteria (blue-green algae) have been identified as one of the promising groups of organisms to produce bioactive metabolites mainly of cytotoxins, enzyme inhibitors, antiparasitic, allelopathic, antimicrobial and AF compounds¹⁵. The exploration of cyanobacterial biodiversity in the marine environment and its associated bioactive metabolites diversity has resulted clinical development of new marine bioactive substances. Till date, several potent compounds from different species isolated of marine cyanobacteria having potent activity against bacteria, fungi, and mycobacterium (Table S1). In these work, antimicrobial activities of cyanobacterial extract was tested on pathogenic bacteria and fungi. Further, the extracts were tested for mammalian toxicity by in *vitro* test (red blood cell toxicity or HepG2 cells). Then various columns chromatographic techniques were used to partially purify the active extracts. Subsequently, thin-layer chromatography (TLC) and gas chromatography-mass spectrometry (GC-MS) were used for active fraction analysis. At last, the potential antimicrobial metabolite generating cyanobacteria was identified and described by 16S rRNA analysis.

Materials and Methods

Collection, isolation, identification, cultivation of cyanobacteria, and extract preparation

Marine cyanobacteria strains were isolated, cultivated in ASN-III medium and identified by using light microscope (Olympus CH20i) at 100X magnification¹⁶. Strains were kept at 27 ± 2 °C, in a light (intensity of 36.45 µmol/m²/s¹; illuminated by cold white fluorescent tubes) and dark cycle of 14 h and 10 h, respectively¹⁷. After the incubation, cells were harvested and extracts were obtained by the cold extraction method. The solvent extracts were centrifuged at 10000 rpm for 10 min and the recovered supernatant was subsequently dried and stored at -20 °C. Thus, obtained crude extracts of the cyanobacterial strains were dissolved in DMSO for antimicrobial assay.

Phylogenetic analysis of 16S rRNA gene and amplification by PCR

The polymerase chain reaction (PCR) was used for amplifying the 16S rRNA gene of the isolated DNA of Geitlerinema carotinosum (Geitler) Anagnostidis CNP 4003. CYA106F (5' CGG ACG GGT GAG TAA CGC GTG A 3') and CYA781 (R)b (5' GAC TAC TGG GGT ATC TAA TCC CAT T 3') primers were used as illustrated earlier by Maruthanayagam et al.¹⁸. The PCR quantities were detained 30 µl having 10 µl of Milli-Q water, one ml of each primer (10 pmol/ml), 16 µl of PCR mixture 2X Master Mix (Prime), and two µl (& 20 ng) of DNA. My gene thermocycler (Applied Biosystems, Foster City, CA) was used for PCR amplification with the following conditions: four min at 94 °C for initial denaturation; 35 cycles of denaturation at 94 °C for four min, annealing at 50 °C for one minute, and extension at 72 °C for 1.3 min; followed by seven minutes of final extension at 72 °C. Agarose gel electrophoresis (1.2 %) and RPC purification kit (Real Biotech Corporation (RBC), Bangiao City, Taiwan) were used to confirm the PCR products (589 bp DNA) and their purity, respectively. Reverse and forward primers were used for sequencing the gene. The BLAST system was used for analyzing the 16S rRNA gene sequences to identify their closest relatives. The public domain GenBank database was used for the gene sequence deposition. ClustalW multiple alignment algorithms were used for multiple alignments and created with reference to the selected GenBank sequences. In the analysis of alignment positions, sequences with one or more gaps or ambiguities were omitted. Neighborjoining method and the bootstrap method of phylogeny with 1500 replications were used for determining the evolutionary history using MEGA 5.0^(ref. 19). The entire place enclosed gaps and missing data were removed.

Antimicrobial screening test

Antimicrobial activities of extracts were screened against the bacterial strains S. aureus ATCC 25923, M. (National Institute of Research in smegmatis India), Pseudomonas Tuberculosis, Chennai, aeruginosa ATCC 27853, Klebsiella pneumoniae ATCC 17106, E. coli ATCC 35218, and a fungal pathogen, C. albicans ATCC 10028. The nutrient agar and Sabouraud dextrose agar were used to maintain the bacterial and fungal cultures, respectively. About 0.85 % NaCl was used for suspending the microbial strains and the turbidity was adjusted to 108 CFU/ml, equivalent to 0.5 MacFarland standards according to NCCLS. Using the non-toxic cotton swab, the cell suspension was inoculated onto agar and incubated for 20 min at 37 °C). Each crude/fractionated extract was estimated mostly by the disc or well diffusion assay according to Kirby-Bauer method²⁰. For the purpose, each those extracts were individually suspended in sterilized DMSO at the concentration of 100 µg or 1 mg and then loaded on to 6-mm Whatman No. 3 filter disc (at a concentration of 100 µg/disc) or well. DMSO was used as a control solvent. Thereafter, those extract spotted plates (were incubated (37 °C, 24 h) and the antimicrobial potential was assessed based on the zone of inhibition (millimeters).

Silica gel column chromatography

The silica gel (100-200 mesh) was prepared and the active fraction was loaded, and separated by the solvent gradient. The collected fractions were spotted in the in house prepared TLC plates (0.25-mm) and kept in the developing chamber (30 % EtoAC: 70 % PE). After developing the plates, 5 % H₂SO₄: methanol solution was sprayed on it. Then the plates were placed in the iodine chamber and heated in hot air oven to remove the excess iodine vapor and finally to observe the spot in UV Transilluminator. As observed, they were similar in composition, as shown by thin-layer chromatography, and combined and dried. The active fraction with antimycobacterial activity against *M. smegmatis* was selected and used for further analysis.

Evaluation of minimal inhibitory concentration by broth macro dilution method

The minimal inhibitory concentration (MIC) was analyzed by using fraction 6 to determine the least concentration of the active fraction that can cause noticeable growth inhibition and bactericidal effects against the M. smegmatis. A fresh colony of M. smegmatis was inoculated in nutrient broth and grown with vigorous aeration for overnight at 37 °C. Ten (10) μ l of the mid-exponential culture (1×10⁶) cells/ml) were mixed with the active fraction dissolved in methanol at concentrations of 50 - 3000ug in one ml total volume of nutrient broth and incubated at 37 °C for 24 h with gentle agitation. After incubation, the culture pellets were gained by centrifugation at 10000 rpm (REMI Cooling Centrifuge 24 BL) and re-suspended in 0.1 ml saline, and the whole suspension was swabbed onto the respective agar plates and allowed to grow for a further 24 – 48 h at 37 °C and then the MIC and MBC were determined²¹.

In vitro cytotoxic assay using marine cyanobacterial extracts

The HepG2 human liver cancer cell line obtained from the National Center for Cell Science (NCCS), Pune, India, was used to estimate the toxicity of cyanobacterial extract by using MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl-2H-tetrazolium bromide) assay²². DMEM medium (Sigma-Aldrich, St. Louis, MO, USA) (10 % fetal bovine serum (Gibco) and 100 U/ml of penicillin and 100 µg/ml of streptomycin as antibiotics (Gibco)) was used for the cultivation of HepG2 cells in 96 well plates at 37 °C in a humidified atmosphere of 5 % CO₂ in a CO₂ incubator (Forma, Thermo Scientific, USA). Cells from passage 15 or less were used for the entire experimentation. Briefly, 96-well flat-bottom tissueculture plates in the complete culture medium for cells $[(5 \times 10^3 \text{ cells } (200 \ \mu \text{/well})]$ seeding. After overnight-seeding, the extracts were added and incubated for 24 hrs in the same CO_2 incubator. DMSO (10 %) added well was used as a positive control. About 20 µl of MTT solution [5 mg/ml in phosphate-buffered saline (PBS)] was added to each well after 24 and 48 h, covered with aluminum foil and incubated for 4 h at 37 °C. After the incubation, 100 µl of DMSO was added to each well to suspend the purple formazan product. The 96-well plate reader (Bio-Rad, Hercules, CA, USA) was used to measure the absorbance at 570 nm (measurement) and 630 nm (reference). Mean of the three replicates were used to calculate the percentage of inhibition as follows: Percentage of Inhibition = (Mean OD of untreated cells (control) - Mean OD of treated cells (treat))/ Mean absorbance of untreated cells (control).

Spector *et al.*²³ were described as the acridine orange and ethidium bromide staining. About 25 μ l of AO and EB solution (3.8 μ M of AO and 2.5 μ M of EB in PBS) was added to the each cell suspension (5×10⁵ cells) and observed under a fluorescent microscope (Carl Zeiss, Germany) with a UV filter (450 - 490 nm). Three hundred cells per sample were counted in triplicate for each dose point. In addition, the morphological alteration was photographed.

Hemolytic Assay

Sodium citrate-EDTA containing centrifuge tubes (15 ml capacity) were used to collect human blood samples. After the collection, blood was transferred to a 50 ml centrifuge tube. To the blood, 30 ml of calcium- and magnesium-free PBS was added and centrifugation at 4000 rpm for 10 min until turned clear and colorless. supernatant The supernatant was discarded and the procedure was repeated twice to obtain clean cells. These cells were kept in PBS (20 ml) and used within 2 weeks. To the 0.5 ml cell suspension in PBS, 5 ml of the test material from a stock solution was added. The number of compounds 93.75, 187.5, 375, 750, 1500, and 3000 (final erythrocyte concentration μg approx. 0.5×10^{9} /ml) were used for testing. The mixture was incubated for one hour at 37 °C with mild shaking and then centrifuged at 4000 rpm for 10 min. The obtained supernatant was diluted 10-fold with PBS and the optical density was measured at 540 nm. PBS or 0.1 % Triton X-100 (w/v in water) was used for calculating the 0 and 100 % lysis and the tests were performed in duplicate²⁴.

Phytochemical analysis

TLC study and developing a solvent system

Precoated silica gel G 60 F254 was purchased from Merck, Germany and the solvents used were of analytical grade. A solution of the active fraction at a concentration of 5 mg/100 μ l was prepared in ethyl acetate. Using a capillary tube, the active fraction was loaded in the Silica gel 60F254 TLC plate and developed with solvent system Acetonitrile: Heptane: Chloroform (1:4.5:4.5 V/V/V) for satisfactory separation of the compounds. After the development of plates, they were air-dried and the numbers of spots were noted. Spots were observed by spraying with various spraying reagents as follows: Ehrlich's reagent (0.1 g of p-dimethyl aminobenzaldehyde in five ml of hydrochloric acid and methanol; plates heated for 20 min at 50 °C) for sulfonamides, amines, indoles, and

ergot alkaloids, Dragendorff's reagent (0.011 g and 0.018 g dissolved in acetic acid and makeup into 10 ml of distilled water) for alkaloids, ninhydrin (0.1 g of ninhydrin dissolved in 9.5 ml of pyridine and 0.5 ml of acetic acid, heated to 110 °C until reddish spots) spray for peptides, phenol (Folin-Ciocalteu) reagent for phenolic compounds, 10 % sodium hydroxide reagent for flavonoids, anisaldehyde-sulphuric acid (0.75 g of anisaldehyde dissolved in 12.5 ml of ethanol and 0.125 ml of concentrated sulphuric acid) for sterols and terpenes, and ferric chloride reagent (5 % in methanol) and potassium hydroxide reagent (10 % in methanol) for Anthraquinone.

GC-MS analysis of active fraction

The coupled GC-MS was used for analyzing the active fraction (1 mg/1 ml ethyl acetate). THERMO GC-TRACE ULTRA VER: 5.0 analyzer with a flame ionization detector was used for the separation of compounds and analysis. DB 35-MS capillary standard non-polar column (length = 30 m, diameter = 0.25 mm, film thickness = $0.25 \mu m$) was used for separation in GC-MS. The carrier gas was helium and the total GC run time was 40.50 min. 70 °C was kept as the initial oven temperature and it was ramped at 10 °C/min until it reached 250 °C. The injector temperature was 250 °C. Ethyl acetate was used as a solvent to separate the compounds in the extracts and the split ratio 1:10. (http://www.sisweb.com/software/ms/nist.htm). The closest match with the highest probability in the NIST Mass Spectral Library was recorded and interpreted.

Results

A total of 12 filamentous forms of marine cyanobacteria (Aphanocapsa littoralis Hansgirg CNP 2022, Phormidium abronema Skuja CNP 2012, Oscillatoria annae van Goor CNP 1020, Anabaena circinalis Var. crassa Ghose CNP 2061, Synechococcus pevalekii Nag. CNP 7001, G. pseudacutissimum Geitler) Anagnostidis CNP 1019, Oscillatoria princeps Vaucher ex Gomont CNP 1014, Anabaena constricta (Szaffer) Geitler CNP 5007, Phormidium valderianum (Delp.) Gomont CNP 4011, Oscillatoria salina Biswas CNP 1006, G. carotinosum (Geitler) Anagnostidis CNP 4003 and O. boryana Bory ex Gomont BDU 91451) belonging to the genera Phormidium, Oscillatoria, Aphanocapsa, Synechococcus, Anabaena, and Geitlerinema were taken for the initial screening. Classical methods based on morphology were used for identifying the strains (Desikachari, 1959) and G. carotinosum CNP 4003 (Fig. 1a) was further confirmed by phylogenetic analysis (Fig. 1c).



0.001

Fig. 1 — Morphology and molecular identification of *G. carotinosum* CNP 4003: (A) Microscopic image of *G. carotinosum* CNP 4003, (B) amplification product of *G. carotinosum* CNP 4003 16S rRNA (*a - G. carotinosum* CNP 4003, *m -100 bp marker*), and (C) phylogenetic tree

Therefore, classification by 16S rRNA gene sequencing was reserved to find out the genus for certain cases. The PCR amplification of the 16S sequence of *G. carotinosum* CNP 4003 (Fig. 1b) resulted in 589 bp DNA and it was used for the BLAST search. The DNA isolated from this cyanobacterium exhibited sequence resemblance with the DNA of another cyanobacterium (accession number AJ621834) and the sequence was deposited in the GenBank database with accession number KC404068.

In this study, organic crude extracts from the above listed cyanobacteria were screened (Table S2) for

their antifungal and antibacterial activity against Candida, Gram-negative and Gram-positive strains of bacteria. Preliminary testing for the antimicrobial study of extracts was performed by the disc diffusion and agar well method. The best antibacterial activity was observed in the extract of G. carotinosum CNP 4003 at 100 µg /disc and 1000 µg/well against S. aureus ATCC 25923, E. coli ATCC 35218, and M. smegmatis with inhibition zones of 6, 8, and 11 mm, correspondingly (Fig. S1). The extract of G. carotinosum CNP 4003 showed the highest inhibition level against M. smegmatis. The cytotoxicity result (Fig. 2) revealed that the potent

extract of *G. carotinosum* CNP 4003 was moderately cytotoxic to mammalian cell line HepG2 (IC₅₀ 175 μ g/ml) with antimycobacterial activity against *M. smegmatis* at 1 mg/well. Furthermore, this extract was fractionated by silica gel chromatography and antimycobacterial activity was checked for 9 pooled fractions of C17 (F1-F9). Of all the tested fractions, fraction 6 showed the maximum inhibition activity at 250 μ g from per disc concentration with 8 mm of inhibition zone (Fig. S2). This fraction was chosen to determine the MIC. Fraction 6 was serially diluted 50 to 3000 μ g and MIC value was determined



Fig. 2 — Cytotoxic activity of active fraction and extract of *G. carotinosum* CNP 4003: Photomicrographs showing the effect on untreated (Control), extract of *G. carotinosum* CNP 4003 ($IC_{50} = 175 \ \mu g/ml$) and active Fraction F6 of *G. carotinosum* CNP 4003 ($IC_{50} = 63 \ \mu g/ml$)



Fig. 3 — Hemolytic activity of active fraction

(Fig. S3). Further, the treated cells were collected and plated for viable count analysis. The results showed that the MBC and MIC of the fraction 6 was 800 and 1500 µg as tested on the *M. smegmatis* strain. Very low hemolytic activity, i.e. 63 % (Fig. 3) was observed at 93.75 µg/ml, whereas the highest hemolysis (95 %) was observed at the concentration of 375 µg/ml. Further, the active fraction F6 was relatively cytotoxic to liver cell line HepG2 (IC₅₀ 63 μ g/ml). Qualitative analysis of the active fraction was given in Table 1 and Figure 4. The results clearly showed that active fraction may consist of indoles or terpenes or peptides. Finally, GC-MS were used for analyzing the chemical composition of the active fraction. A high percentage of the probability for five chemical compounds was noticed in the active fraction with peaks at 3.29, 7.98, 10.10, 12.69, 15.60, and 30.44 RT. In the commercially available databases, the spectra of twenty-five peaks did not match, signifying the newness of these molecules (Table 2).

Discussion

The purpose of this study was to identify the active cyanobacterium among the marine isolates against pathogenic microbes and to select potential candidate for future research. In a similar study against the ethyl acetate extract of Mycobacterium, Aerococcus sp. was found to be active against M. smegematis with an inhibition zone of 8 ± 0.1 mm²⁵. Mariita et al.²⁶ reported antimycobacterial activity of MeOH extract of Entada abysinnica (2.0 and 1.0 mg/ml) showed antimycobacterial activity against М. tuberculosis, M. fortuitum, and M. smegmatis. The cell-free extract of Nostoc muscorum Ag (U2301) and Fischerella ambiguia (Näg) Gom (U 1903) exhibited significant antimicrobial

Table 1 — Constituent analysis of active fraction ofG. carotinosum CNP 4003				
Chemical Constituent (class)	Indicating Reagent	Results		
Sulfonamides, amines, indoles, and ergot alkaloids	Ehrlich's reagent	+		
Alkaloids	Dragendorff's reagent	_		
Peptides	Ninhydrin	+		
Phenolics	Phenol reagent	+		
Flavonoids	10% NaOH	_		
Sterol	Phosphomolybdate	+		
Anthraquinone	10% sodium hydroxide	_		
Tannins	Ferric chloride reagent	_		
Terpenes	Anisaldehyde- perchlorate	+		
+ indicates presence, - indicates	absence			



Fig. 4 — Qualitative identification of compound classes in the active fraction of *G. carotinosum CNP 4003*: Extract was loaded on silica gel plates and developed with acetonitrile:heptane:chloroform (1:4.5:4.5 v/v/v) solvent mixture. After development, the plate was observed directly (Panel A) or under UV light at 254 nm (Panel B) or 366 nm (Panel C). Other plates were sprayed (and developed subsequently as outlined in Methods) with Anisaldehyde–H₂SO₄ (Panel D), 10 % KOH (Panel E), Ehrlich's (Panel F), ninhydrin (Panel G), Folin–Ciocalteu (Panel H), Ferric chloride (Panel I), anisaldehyde-HClO₄ (Panel J), 10 % NaOH (Panel K) and Dragendorff's reagent (Panel L). Red arrow indicates presence of indoles (purple spot). Green arrow indicates presence of peptides (pink spot). Blue arrow indicates presence of terpenes (dark blue spot).

Table 2 — Composition of active fraction of <i>G. carotinosum</i> CNP 4003 as revealed by gas chromatography mass spectrometry (GC-MS)			
Peak retention time (min)	% of area	Close match	% of similarity
3.29	75.66	Acetic acid, ethyl ester	93.50
5.30	3.29	UI	-
7.98	0.30	1,3-Bis(4-chlorobenzyl)-5,6-dihydrobenzo[f]quinazoline	53.44
10.10	0.32	Cycloheptasiloxane, tetradecamethyl	97.50
12.69	0.23	Hexadecamethylcyclooctasiloxane	94.52
14.54	0.13	UI	-
14.99	0.41	UI	-
15.60	0.18	Cyclononasiloxane, octadecamethyl	83.79
18.35	0.25	UI	-
18.81	0.27	UI	-
19.81	3.07	Neophytadiene	38.91
20.44	0.85	UI	-
20.81	1.77	3,7,11,15-Tetramethyl-2-hexadecen-1-ol	35.21
22.88	0.41	UI	-
23.32	0.11	UI	-
23.95	8.63	Hexadecanoic acid	36.86
25.05	0.21	UI	-
26.12	0.56	UI	-
26.70	0.23	UI	-
27.98	0.17	UI	-
29.42	0.12	UI	-
29.95	0.18	UI	-
30.44	1.15	5,10-Dihexyl-5,10-diihydroindolo[3,2-b]indole-2,7-dicarbaldehyde	58.52
31.20	0.13	UI	-
32.44	0.16	UI	-
33.03	0.34	UI	-
33.73	0.11	UI	-
34.44	0.34	UI	-
38.23	0.15	UI	-
40.25	0.27	UI	

The data represents the relative area of the peaks in %. Peaks were identified by the NIST Mass Spectral library. Compounds that have a probability lower than 75 % to the closest match with the library are marked as unidentified (UI).

activity against *M. tuberculosis*²⁷. Extract of the cultured cyanobacterium Eucapsis sp. (UTEX 1519) exhibited the most potent activity (MIC 9.7 µg/ml) against *M. tuberculosis*²⁸. Malyngolide isolated from L. majuscula showed potential antibacterial activity against *M.* smegmatis²⁹. The active fraction of Hapalosiphon sp. displayed a minimum inhibitory concentration at 125-2000 μ g/ml against strains of *M*. Tuberculosis³⁰. Isolation of fischambiguine B from the cultured cyanobacterium, Fischerella ambigua (UTEX 1903) showed activity against M. tuberculosis with a MIC value of 2 μ M and did not exhibit cytotoxicity on Vero cell line³¹. Brunsvicamide B from Tychonema sp. isolated from the pond of a sugar factory potentially inhibited M. tuberculosis enzyme MptpB with an IC₅₀ value of 7.3 μ M³². Also Fischerella sp. isolated from Neem tree bark exhibited antimycobacterial activity against $M. tuberculosis^{33}.$

Among the 12 strains of cyanobacteria tested for antimicrobial activity, G. carotinosum CNP 4003 showed better antimicrobial activity than the other strains. Extract of G. carotinosum CNP 4003 showed distinct antimicrobial activity against S. aureus ATCC 25923, E. coli ATCC 35218, and M. smegmatis. Results of the hemolytic and cytotoxic activity of extract. and active fraction against human erythrocytes and HepG2 cell line showed its viability as a future drug candidate particularly for internal use. Bioactive molecules from marine cyanobacteria are well-known for their toxic effects against animals and some of the eukaryotic microbes. It is also reported that these structurally diverse antimicrobial compounds produced by cyanobacteria are discrete substances and differ from the cyanotoxins. Considering these properties, we have evaluated the cytotoxicity of the antimicrobial active extract against human erythrocytes and HepG2 cancer cells by hemolytic and MTT assay. This result demonstrated the that active fraction showed good antimycobacterial activity and high toxicity to human red blood cells and cancer cell lines. The assay indicated that the presence of toxic activity could be understood to the presence of cytotoxic activity for the active fraction.

Compounds from cyanobacteria have been accounted for having different modes of antibacterial actions. Assessment of cyanobacteria peculiarly growing on black band disease of corals revealed the production of antibacterial compounds in them and also indicated that their activity depends mainly on ecological conditions³⁴. Preliminary screening of organic and aqueous extracts of cyanobacteria, Anabaena strains, isolated from terrestrial environment exhibited antibacterial and cytotoxic activity³⁵. Asthana et al.³⁶ explored biochemical and antibacterial properties of Antarctic strain, Nostoc sp., and stated that antibacterial activity was specially connected with the habitat. Extracts from evanobacterial mats of hot springs inhibited the growth of Klebsiella pneumoniae, Salmonella enteric, Micrococcus luteus, Shigella sonnei, Bacillus sp., Amphora coffeaeformis and QS of the reporter strain Chromobacterium violaceum CV017 and Agrobacterium tumefaciens NTL437. Pramanik et al.38 reported the antimicrobial activity of cyanobacteria isolated from Sundarbans mangrove forest against P. aeruginosa, E. coli, B. subtilis, and S. aureus multiple drug-resistant clinical isolates (MIC 0.25-0.5 mg/ml). Furthermore, these extracts were shown to be nontoxic to human colon adenocarcinoma cell line at the same MIC value. Biomass extract of antibacterial activity of Geitlerinema sp. was recorded by Caicedo et al.³⁹ and they further reported 100 % growth inhibition of Bacillus subtilis by the crude extract eluted with MeOH and isopropanol.

Jaiswal et al.40 and Kim & Kim41 reported the antifungal activity of extracts from Microcystis aeruginosa and Nostoc commune FA-103 against the growth of phytopathogenic fungus, Fusarium oxysporum f. sp. lycopersici and Rhizoctonia solani. Another study demonstrated the antimicrobial potency of Nostoc strain ATCC 53789 against Sclerotinia sclerotiorum, Rosellinia sp, Penicillium expansum, and Armillaria sp at 0.25 g/ 1^{42} . In an investigation by Pawar & Puranik⁴³, methanolic extract of Oscillatoria limosa NMU-31 displayed a good antifungal activity against Aspergillus flavus with an inhibition zone of 28 mm diameter. Extracts obtained from Geitlerinema strain Flo1 by using Amberlite XAD-1180 resin exhibited against marine yeast, Rhodosporidium activity sphaerocarpum⁴⁴. All these studies exposed the hidden novel antimycobacterial pharmacophores in marine cyanobacteria. Thus, it's understood that the primary extracts of various species of marine cyanobacteria displayed promising antimycobacterial activity against Mycobacterium smegmatis. The purification of these crude extracts by different chromatographic techniques and structure elucidation by various spectroscopic analyses, such as IR, NMR, and MS is in progress.

Conclusion

The results of the present study justified the promising antimycobacterial activity of the tested cyanobacterial crude extract. Further research is required to isolate, and identify the active principle(s) from cyanobacteria, and to evaluate their mode of action(s). The entire work specified at this point emphasize that the isolated cyanobacteria from the marine environment can be a potent antimicrobial agent.

Supplementary Data

Supplementary data associated with this article is available in the electronic form at http://nopr.niscair.res.in/jinfo/ijms/IJMS_49(07)1165-1174 SupplData.pdf

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Conflict of Interest

The authors report no conflicts of interest. The authors alone are responsible for the content and writing of this article.

Author Contributions

MS: Conceptualization, Project administration, Writing - review & editing; VM: Formal analysis, Investigation, Methodology, and Writing - original draft; MN: Formal analysis, Investigation, and Methodology.

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