Isolation, Phylogenetic Analysis of the Microbial Community Associated with the Red Sea Sponge *Ircinia Echinata* and Biological Evaluation of their Secondary Metabolites

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Abstract

The present study was undertaken to provide information about bioactive compounds of the microbial community associated with *Ircinia echinata* marine sponge collected from the Red Sea. This work investigated the antioxidant and cytotoxicity properties of the tested extracted crude extract. Sponge was taxonomically identified as *Ircinia echinata*. Moreover, taxonomic and phylogenetic analyses of the bacteria on the basis of 16S rRNA gene sequences showed that dominant phylum was proteobacteria. The results of HPLC analysis revealed that the bacterial crude extracts exhibited highly significant quantities of total phenolic contents in crude extracts of the two tested bacterial isolates. The highest level of total phenolic contents was measured in the extract was shown in extracts HA-MS-41 (3.238±0.03 mg gallic equivalent/ml). Whereas, the lowest value was shown in extract HA-MS-118 (2.946±0.10 mg gallic equivalent/ml).

The results of the current study cleared out that extract HA-MS-41[JX484800] contained gallic and tannic acids (1.084 and 1.413 mg/ml), whereas extract HA-MS-118[JX484801] contained gallic (1.005 mg/ml) but didn't contain tannic acid. The highest level of total flavonoid content was measured in the extract HA-MS-41 extract (5±0.23 mg catechin equivalent / ml). Whereas, much lower value was observed in the HA-MS-118 extract (2.5±0.26 mg catechin equivalent / ml). Carotenoid compounds significantly varied among the two bacterial extracts; the highest total carotenoid concentration was observed in the extract HA-MS-41 (1.165µg/ml). Whereas, lower values were observed in the extract HA-MS-118 (0.185 µg/ml). In the meantime, GC/MS profiling for the ethyl acetate extract of the studied sponge-associated bacteria revealed the presence of various phytochemical compounds having different biological activities. Results of FTIR spectroscopic studies have revealed the presence of various chemical constituents in ethyl acetate extract of the HA-MS-41 and HA-MS-118 bacterial isolates.

The results of the elemental analysis indicated that the bacterial extracts contained major elements like Na, Mg, Al, Si, S, K, Ca, P and Cl, besides the major elemental constituents in all living organisms; C, N and O. The results showed considerable concentrations of many halides (Fe, Cu, Zn and Br). DPPH scavenging activity in terms of IC50 values was shown to be high in case of the extract HA-MS-41 (1.045 mg/ml) and in case of the extract HA-MS-118 (1.450 mg/ml). IC50 values of the antioxidant activity of bacterial extracts using NO assay for the tested two extracts were 5.98 and 6.222 %. Moreover, potent antioxidant activity using ABTS+ assay with percentages inhibition of IC50 = 3.404 and 1.64 mg/ml for HA-MS-41 and HA-MS-118, respectively. In the present study highly significant total antioxidant capacity values was also obtained by using phosphomolybdenum assay in the bacterial extract suggesting the presence of effective antioxidants in the bacterial extract.

The high potency of these isolates is mainly attributed to the presence of various classes of potentially active secondary metabolites detected in them. Indeed phenols, tannins, and alkaloids identified in the tested extracts have been reported to possess antioxidant and anticancer activities. The relative wide range of these biologically properties may results from the individual or from the combined modes of action of compounds belonging to the identified groups of constituents. The cytotoxicity screening studies revealed that the maximum cell growth inhibitory effects of bacterial crude extracts against the tested cancer cell lines; HepG2; HCT and MCF-7 were found to be 22.7, 14.6, and 12.2 µg/ml, respectively for HA-MS-41, and 36.5, 23.6, and 11.9 µg/ml, respectively for HA-MS-118. These results were confirmed by a protein kinase inhibition specialized screening revealing an inhibition activity of 89.3±2.1 and 94.3±1.15 % for HA-MS-41 and HA-MS-118.

**Keywords:** Sponge Associated Bacteria; Phylogenetic Analysis; Bioactive Metabolites

**Abbreviations:** NIWA: National Institute of Water and Atmospheric Research; PCR: Polymerase Chain Reaction; SEM: Electron Microscope; EDX: X-ray Spectrophotometer; DPPH: Diphenyl–α-picrylhydrazyl; RP: Reversed-Phase; BHT: Butylated Hydroxytoluene; BHA: Butylated Hydroxyanisole; TAO: Total Antioxidant Capacity; HepG2: Hepatocellular Carcinoma
Introduction

The marine life has an immense potential as a source of novel bioactive compounds which has been barely investigated. Natural products from marine sponges exhibit unique structural features which are of paramount interest for the scientists for the identification of new lead structures for potential drugs. Sponges (phylum Porifera) are known to be extremely rich sources of bioactive compounds, mainly of secondary metabolites of pharmaceutical and medical relevance. The development of secondary metabolites from the Red Sea sponges offer a potential for producing novel drugs and prototypes and has been a subject of intense research due to their immense bioactive nature [1-9]. They are also well known to be hosts for a large community of microorganisms [1,10]. Associations between higher organisms and microorganisms often occur as mutualistic or symbiotic relationships, by providing benefits for both in terms of protection and nutritional requirements [11-13]. These microorganisms can make up 35–60% of the total sponge mass and are responsible for the production of numerous bioactive compounds with unique, complex and different structure compared with terrestrial microorganisms [4,10]. Production of secondary metabolite is influenced by the environment. Inhibitory interactions among sponge-associated bacteria that occur within the sponge surface are of great interest to search for secondary metabolite-producing bacteria. In the meantime, they are the richest source of microbial community. So far, the bacterial community associated with sponge in the Red Sea remains unexplored. They produce diverse and unique bioactive secondary metabolites. This is an adaptation strategy to thrive in the extreme environmental conditions of the sea and as a defense strategy to escape from predators by the marine invertebrates especially soft bodied animals like sponges [14].

Since the marine environment is composed of a myriad of ecologically distinct habitats, it is not surprising that bacteria adapted to life in the marine environment are both diverse, phylogenetically distinct, and maintain survival adaptations that differ from bacteria that occur on land. These bacteria are widely reported to be highly potential as the producer of antimicrobial bioactive compound [11,12,15]. Numerous types of infectious diseases caused by pathogenic bacteria or yeast are one of basic problems in biomedical field that need to be seriously controlled. Therefore, the isolation of the microorganisms responsible for the origin of the bioactive products of interest will enable large-scale production of marine pharmaceuticals by means of fermentation technology. Despite having more than 2250 km coastline along the Red Sea, Egyptian studies concerning marine natural products related to drugs discovery are still very scarce. So far only a few classes of Egyptian marine organisms have been investigated for their chemical and pharmacological properties. Therefore it is believed that the identification of Egyptian organisms with significant biotechnological potential is an important tool for the discovery of new drugs. The main objective of the present study is to isolate and carry out a phylogenetic analysis of the microbial community associated with the Red Sea Sponge *Ircinia echinata* in order to explore their different secondary metabolites of biological activity and to characterize their potential as antioxidant and anticancer.

Materials and Methods

Sponge Collection

Sponge samples were collected in May, 2010 from the Red Sea by SCUBA diving in (Figure 1). The physical, chemical, and biological characteristics of the Red Sea marine environment have been very recently extensively investigated [1-9; 16-31]. Samples were cut from the sponge with a dive knife while wearing latex gloves and individual pieces were transferred to separate plastic sample collection bags, brought to the surface, maintained at ambient seawater temperature and transported to the laboratory in the same day of collection. Sponge specimens were rinsed in sterile seawater, cut into pieces of ca. 1 cm$^3$, and then thoroughly homogenized in a sterile mortar with 10 volumes of sterile seawater. The supernatant was diluted in ten-fold series (10$^{-1}$, 10$^{-2}$, 10$^{-3}$) and subsequently plated out on agar plates.

![Figure 1](image_url)

Figure 1: Map of Egypt showing (A) the sampling sites at Hurghada and (B) the Red Sea Ircinia echinata Sponge.
**Taxonomic Identification and Processing of the Sponge Sample:** Sponge was taxonomically identified by Prof. Michele Kelly at National Institute of Water and Atmospheric Research (NIWA) Ltd., Auckland, New Zealand as *Ircinia echinata* (Order: Dictyoceratida; Family: Irciniidae). A voucher specimen has been deposited at the Auckland War Memorial Museum. A voucher specimen of the sponge sample has been deposited in the NIWA Museum, under registration number NIWAKD 6672. Sponge was transferred to plastic bags containing sea-water and transported to the laboratory. Sponge specimen was rinsed in sterile seawater. Fresh specimen of each sponge was used for isolation of the sponge-associated bacteria. A section of sponge was immediately used to culture the sponge associated bacteria. Another section of sponge was immediately frozen on dry ice and stored at -80°C. The sponge tissue was then freeze-dried prior to DNA extraction.

**Isolation of Sponge Associated Bacteria:** Two growth media were prepared to isolate a wide range of the sponge associated bacteria; Marine Agar [32], and ISP medium 2 [33] served as general rich media to grow many heterotrophic marine bacteria. All media were supplemented with 0.2 μm pore size filtered cycloheximide (100 μg/mL), nystatin (25 μg/mL) and nalidixic acid (25 μg/mL). Cycloheximide and nystatin inhibit fungal growth, while nalidixic acid inhibits many fast-growing Gram-negative bacteria [34]. All media contained Difco Bacto agar (18 g/L) and were prepared in 1 L artificial sea water (NaCl 234.7 g, MgCl2·6H2O 106.4 g, Na2SO4 39.2 g, CaCl2 11.0 g, NaHCO3 1.92 g, KCl 0.64 g, KBr 0.96 g, H3BO3 0.26 g, SrCl2·0.24g, NaF 0.03 g and ddH2O to 10.0 L) [35]. The inoculated plates were incubated at 28°C for 5 - 10 days. Distinct colony morpho-Types were picked and re-streaked until visually free of contaminants. Isolates were inoculated into liquid media ISP medium 2. The isolates were maintained on plates for short-term storage and long-term strain collections were set up in medium supplemented with 30% glycerol at -80°C [36].

**Extract Preparation and Cytotoxic Activity Screening:** Bacterial isolates were cultured in 500 mL Erlenmeyer flasks containing 200 mL of ISP2 medium. The liquid cultures were grown for 7 - 14 days depending on their growth rate at 30°C while shaking at 150 rpm. Crude extracts were prepared from whole cultures; containing cells and broths by ethyl acetate equal volume/three times, evaporated, lyophilized and dissolved in 1:1 v/v DMSO/H2O and kept at 4°C until use for cytotoxic assay. The MTT [(3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide)] assay which Measuring the mitochondrial re-ductase enzyme activity of viable cells that Could reduce MTT to formazan, (bromide) assay which Measuring the mitochondrial re-ductase activity of viable cells that Could reduce MTT to formazan, giving a purple color, was used for screening the cytotoxicity of the bacterial extracts [37]. The cytotoxic activity of the microbial crude extracts was determined against three established cancer cell lines; MCF-7, HepG2 and HCT cells. Based on the values of the IC50, the most potent four bacterial extracts which exhibited the lowest IC50 were chosen for further study of their phylogenetic analysis and identification.

**Molecular Identification of the Bacterial Isolates**

**Genomic DNA Extraction:** Bacterial DNA was extracted using QIAgen DNAeasy Blood and Tissue Kit (Cat. # 69506), following manufacturer’s guidelines

**Polymerase Chain Reaction (PCR) Amplification of 16S rRNA:** Following extraction of genomic DNA, polymerase chain reaction (PCR) was carried out in order to amplify the 16S rRNA gene, the primers used to amplify the 16S rRNA gene were two universal bacterial primers: Forward primer 27f (5’-AGAGTTTGATCTCGTATCAAGAG-3’) and reverse primer 1492r (5’-GGTTACCTTGTTACGACTT-3’). The PCR conditions used were the same as those described previously [39]. The PCR products were purified and sequenced by the GATC-Biotech. Company (Tübingen, Germany). The sequences were compared with known sequences in the GenBank nucleotide database and the species level was identified as the nearest phylogenetic neighbor with 98% - 100% similarity [40]. Sequence alignment and phylogenetic analysis were performed using the Mega-5 software package [41]. Tree construction was conducted using neighbor joining algorithm (Jukes-Cantor correction) with bootstrap values based on 1000 replications. The 16S rRNA gene sequences the isolates were deposited in GenBank under the accession Numbers indicated in brackets: HA-41 [JQ768238] and HAMS- 118 [JQ768240].

**Chemical Investigations of the Bacterial Isolates**

**Analytical HPLC analysis:** 20 μl sample extract was analyzed with a Exclipse XDB C18 (5 μm, 4.6 *150 mm) column using a mobile phase consisting 1% (v/v) formic acid in aqueous solution: acetonitrile: 2-propanol (70:22:8), pH 2.5; flow rate: 0.75 ml/min, temperature: 30°C, UV detection at 320 nm; Agilent technologies 1200 series. Phenolic compound concentrations were calculated from the standard curves of the supplied phenolic standards exist in the data library.

**Gas Chromatography- Mass Spectrophotometric (GC-MS) analysis:** Analysis was conducted using Gas Chromatography-Mass Spectrophotometer; GC-MS (Trace DSQ11 MS). MS conditions were as follows: Detector mass spectrometer voltage 70eV and its source temperature was 300°C. The injector temperature was 240°C and the split less mode 0.5μl injection. The HP 55% dimethyl-95% dimethyl tertiary-butyl non-polar column was performed with length 30 cm x 0.25 mm, coating thickness film 0.25 μm. The oven was adjusted at 80°C for 1 min and initial time 1.5 min with 40°C which ended by a final temperature of 300°C and 4 min hold time where the total run time was 15 min. The compounds were identified by comparison with the standards, and also matched with the in-built libraries. The components were identified by comparing their retention times with those of authentic samples, as well as by comparing their mass spectra with those of Wiley 275 Library [42]. Quantitative data were obtained by the peak normalization technique using integrated FID response.

**Infrared Analysis (FT-IR):** Using Perkin-ELMER1430 infrared spectrophotometer, the molecular structure of the extracts
was partial identified. Separated spots of each compound were collected by scratching and dissolved in 1ml ethyl acetate and lyophilized then small discs made from the mixture of about 1mg of the lyophilized material and 300mg of pure and dry KBr were subjected to IR-spectrocope. The measurement were carried out at infrared spectra between 400-4000nm.

**Elemental Analysis:** The bacterial extracts were subjected to the elemental analysis using Scanning Electron Microscope (SEM) with an energy dispersive x-ray spectrophotometer (EDX). This was carried out at the Regional Center for Mycology and Biotechnology, Al-Azhar University, Egypt.

**Determination of Total Phenolic Contents:** Total phenolic compounds in the fungal extracts were determined by the method of [43]. One hundred microliters Folin-Ciocalteau reagent (50%, v/v) in distilled water and 2 ml sodium carbonate (2%, v/v) in distilled water were added and mixed well with 100 µl of fungal extract. After 2 h incubation at RT, the absorbance of the resulting blue color solution was measured at 750 nm. Different concentrations (0 – 0.9 mg/ml) of standard gallic acid were used and processed similarly as sample. The concentration of total phenolic in bacterial extracts was determined as mg gallic acid equivalent in 1 ml of the extract using the standard curve of the gallic acid.

**Determination of Total Flavonoid Contents:** Total flavonoid content was determined by a colorimetric method of [44]. A 0.25 ml of fungal extract or (+)-catechin standard solution was mixed with 1.25 ml of distilled water, followed by the addition of 0.75 ml of 5% sodium nitrite solution. After 6 min, 0.15 ml of 10% aluminum chloride solution was added and the mixture was made up to 2.5 ml with distilled water and well mixed. The absorbance was measured immediately at 510 nm using a spectrophotometer. The results were expressed as mean ml of (+)-catechin equivalents.

a) Bioactivity screening of the bacterial extracts  
b) Anticancer activity screening  
c) Cytotoxicity against cancer cell lines  
d) Cytotoxicity evaluation using viability assay

For cytotoxicity assay, the cells were seeded in 96-well plate at a cell concentration of 1×104 cells per well in 100µl of growth medium. All experiments were carried out in the triplicate. The cell cytotoxic effect of each tested compound was calculated [45,46].

**Protein kinase Inhibitory Effect:** Protein kinase inhibitory effect of the tested bacterial extract was determined by Universal Tyrosine Kinase Kit; Takara (CAT # MK410) [http://www.takara.co.kr/file/manual/pdf/MK410_e.pdf] following the manufacturer’s instructions with some modifications.

In vitro Antioxidant Capacity Screening

**Diphenyl-α-picrylhydrazyl (DPPH) radical scavenging assay**

DPPH radical scavenging assay (Figure 24) of the extracts was performed by using modified previously established methodology by Blois [47], and Amarowicz et al. [48]. Assays were performed in flat bottom polystyrene 96 well microtiter plates. To 100 µL of each sample (1-6mg/ml) in EtOH 25 µL DPPH (1mM) in ethanol was added. The resultant mixture was briefly shaken and maintained at room temperature, in the dark for 30 min. At the end of this period the absorbance (A) of the mixture was measured at 490 nm, using Microplate reader spectrophotometer. Scavenging ratio of DPPH assay was calculated as follows:

\[
\%\text{scavenging} = \left[\frac{(A \text{ control} - A \text{ sample})}{A \text{ control}}\right] \times 100
\]

**Nitric Oxide Scavenging Activity**

The extent of inhibition of nitric oxide radical generation in vitro was followed as per the method reported by Green et al. [49].

**ABTS+ Scavenging Effects**

The antioxidant effect of the leaf extracts was studied using ABTS (2,2′-azino-bis(3-ethylbenzthiazoline-6-sulphonic acid) radical cation decolourisation assay according to the method of Shirwaikar et al. [50].

\[
\text{Inhibition} = \frac{(\text{Control} - \text{test})}{\text{Control}} \times 100
\]

All Measurements were expressed as Trolox equivalents. Standard regression line for Trolox was used to calculate TEAC values.

**Phosphomolybdenum Assay**

This assay is a spectroscopic method for the quantitative determination of antioxidant capacity, through the formation of phosphomolybdenum complex.

**Statistical Analysis**

All experiments were conducted at least with 3 replicates and the results were expressed as mean of 3 independent experiments + SEM calculated using Graph pad prism version 6.0.

**Results and Discussion**

**Recovery of Bacterial Isolates from the Culture Media**

Based on the morphological characteristics and uniqueness, two isolates from sponge Hyrtios erectus. The bacterial isolates were given codes prior to identification (Table 1).

**Molecular Identification of the Bacterial Isolate**

**DNA Base Composition and Sequencing of the 16S Ribosomal DNA (rDNA):** The 16 S rRNA genes were amplified by using general bacterial primers. The forward primer (27 f 5′-CCAGAGTTTGATCCTGGCTCAG-3′) and reverse primer (1492 r 5′-TAC CTT GTT ACG ACT T-3′) were used for amplification of 16S rRNA genes the studied bacterial isolate. The amplified 16S rRNA gene was 1.5 kb.

**Sequence Alignments and Phylogenetic Inference:** The bacterial isolate was identified using a molecular approach. The results showed that the 16S rRNA genes belonged mainly to the phyla proteobacteria. The sequence were deposited in the GenBank database. Accession numbers [JX484800] corresponded to the
strain HA-MS-41 and [JX484801] corresponded to the strain HA-MS-118 (Table 1). For the *Ircinia echinata* sponge sample, bacterial PCR product was sequenced and phylogenetically analyzed (Figures 2 & 3). The present study revealed that the identified bacterial species was Proteobacteria (especially Gamma-Proteobacteria). The phylum Proteobacteria are always found in different sponges from the same or different geographic location. Proteobacteria have been suggested to have varied effects on sponge hosts such as nitrogen fixation and manipulating host reproduction. It has been reported that Proteobacteria produce low molecular-weight biological active compounds with bioactive properties. Proteobacteria were also found to produce enzymes at high levels for degrading protein and polysaccharides (Thomas et al.). Based on our results, phylum Proteobacteria are the predominant bacteria in the *Ircinia echinata* Red Sea sponge. So far different groups of antimicrobial bacteria including α, β and γ proteobacteria have been isolated from different sponges from geographically different regions [51].

**Table 1:** Morphological characteristics and code numbers of the isolated bacteria from the studied sponge samples.

| Sponge        | Isolate code | Colony morphology                                      |
|---------------|--------------|--------------------------------------------------------|
| *Ircinia echinata* | HA-MS-41     | Yellow, creamy and gummy, raised, irregular edge       |
| HA-MS-41      |              |                                                        |
| HA-MS-118     |              | White, creamy, raised, irregular edge                  |

| Bacterial isolate [Accession no.]* | Blastn results (accession number), Max. identity% | Phylum       |
|-----------------------------------|---------------------------------------------------|--------------|
| HA-MS-41 [JX484800]              | *Cedecea davisae* strain DSM 4568, 16S ribosomal RNA, partial sequence (NR_025243.1), 99% | Proteobacteria |
| HA-MS-118 [JX484801]             | *Providencia vermicola* 16S ribosomal RNA, complete sequence (NR_042415.1), 98% | Proteobacteria |

*Accession numbers obtained for the bacterial isolates upon submission in the Gen Bank.

**Figure 2:** Molecular Phylogenetic analysis of the bacterial isolate HA-MS-41 by Maximum Likelihood method. The scale bar indicates 0.5 substitutions per nucleotide position.

**Figure 3:** Molecular Phylogenetic analysis of the bacterial isolate HA-MS-118 by Maximum Likelihood method. The scale bar indicates 5 substitutions per nucleotide position.
Chemical Investigation of the Bacterial Extract: Ethyl acetate extract of the bacterial isolate was subjected to chemical profiling and characterization using HPLC, GC/MS, FT-IR and elemental analysis techniques.

Analytical HPLC Analysis: The results obtained from HPLC analysis against a wide range of phytochemical standards revealed that there were considerable amounts of well-established bioactive compounds such as phenolic compounds, terpenoids and carotenoid compounds.

Phenolic Compounds

The obtained results from the analytical determination of reversed-phase (RP) HPLC against wide range of phytochemical standards revealed that the bacterial exhibited variable patterns of phenolics compounds (Table 2). Chlorogenic acid, 3,5-Dicaffeoyl quinic acid, and 4,5-Dicaffeoyl quinic acid were the main phenolic compounds present in the HA-MS-41 [JX484800] crude extract, whereas Chlorogenic acid and 3,5-Dicaffeoyl quinic acid were present in the HA-MS-118 [JX484801] crude extract. Two of the most powerful phenolic compounds are gallic and tannic acids were also determined by the HPLC analysis for the tested bacterial extracts (Figure 4). The results of the current study cleared out that extract HA-MS-41 [JX484800] contained gallic and tannic acids, whereas extract HA-MS-118 [JX484801] contained gallic acid but didn’t contain tannic acid (Table 3). Gallic acid is commonly used in the pharmaceutical industry. It seems to have anti-fungal, antiviral and antioxidant properties as it helps to protect human cells against oxidative damage. It was found also to show cytotoxicity against cancer cells, without harming healthy cells. Gallic acid is also used to treat albuminuria and diabetes [52].

Table 2: Phenolic compounds concentrations of the tested bacterial extracts.

| Extract            | Phenolic contents (µg/ml) | Chlorogenic acid | Caffeic acid | 4,5-Dicaffeoyl quinic acid | Total |
|--------------------|--------------------------|------------------|--------------|---------------------------|-------|
| HA-MS-41 [JX484800]| 1.865                    | 1.165            | 1.961        | 4.991                     |
| HA-MS-118 [JX484801]| 2.396                    | 5.642            | 2.331        | 10.369                    |

N.D.: not detected. All analyses were mean of triplicate measurements.

Table 3: Concentrations of Gallic and Tannic acids in the tested bacterial extracts.

| Extract            | Concentration of Gallic acid (mg/ml) | Concentration of Tannic acid (mg/ml) |
|--------------------|--------------------------------------|-------------------------------------|
| HA-MS-41 [JX484800]| 1.084                                 | 1.005                               |
| HA-MS-118 [JX484801]| 1.413                                 | N.D.                                |

*ND: not detected. All analyses were mean of triplicate measurements.

Figure 4: Structure of Gallic acid (A) and Tannic acid (B).

Tannic acid, on the other hand, is another phenolic compound that has been found in one of the two extracts. It is a specific commercial form of tannin, a type of polyphenol. Tannins are natural products have large amounts of phenolic rings in the structure and classified into two groups: hydrolyzable and condensed. Condensed tannins contain flavonoids units with several degrees of condensation. Hydrolyzable tannins are considered a mixture of simple phenols with ester linkages in its structure. There are many factors such as alkali compounds, mineral acids, and enzymes that have the ability to hydrolyze tannins [53]. Tannic acid was used as a treatment for many toxic substances. Moreover, it has been stated that the broad spectrum biological activity of some sponges are mainly due to the presence of terpenes, alkaloids and tannins detected in the bioactive fractions [54].

Total Phenolic Contents: Phenolic compounds have been shown to possess important biological activities based on their structural characteristics [1-9]. There are concerns about using synthetic phenolic antioxidants such as butylated hydroxytoluene...
(BHT) and butylated hydroxyanisole (BHA) as food additives because of the reported negative effects on human health [52]. reported also that ethyl acetate fraction exhibited effective antioxidant and antiproliferative activities. The phenolic compounds identified in ethyl acetate fraction could be responsible for the activities. Highly significant quantities of total phenolic contents were found in crude extract of the two tested bacterial isolates equivalents of gallic acid (Table 4). The highest level of total phenolic contents was shown in extracts HA-MS-41 (3.238±0.03 mg gallic equivalent/ml). Whereas, the lowest value was shown in extract HA-MS-118 (2.946±0.10 mg gallic equivalent/ml).

**Table 4:** Total phenolic contents in the bacterial crude extracts expressed as mg gallic acid equivalent/ml.

| Extract            | Concentration of phenolic compounds (mg gallic acid equivalent / ml)* |
|--------------------|---------------------------------------------------------------------|
| HA-MS-41 [JX484800] | 3.238±0.03                                                           |
| HA-MS-118 [JX484801] | 2.946±0.10                                                           |

*Results = Mean (n=3) ±S.E.

**Carotenoids Compounds and Retinol:** Carotenoid contents tested in the bacterial extract were Xanthophyll, β-cryptoxanthin and β-carotene (Figure 5). Xanthophyll, beta-cryptoxanthin, beta-carotene (Table 5) were the major carotenoid compounds investigated in the present study. This group of compounds were reported to have strong antioxidant activity and biochemical properties influencing signaling pathways. Carotenoids was reported to prevent cancers and cardiovascular disorders. Besides, they may protect light-exposed tissues. Ample evidence is available in the literature suggesting carotenoids could possibly lower the risk of stroke, heart diseases, aging, vascular and other metabolic diseases [55]. The antioxidant potential of carotenoids is of particular significance to human health, due to the fact that losing antioxidant-reactive oxygen species balance results in "oxidative stress", a critical factor of the pathogenic processes of various chronic disorders. Carotenoids play a prominent role in protecting bodily cells and thereby act as powerful antioxidants. Moreover, they are proved to be vital for maintenance of proper eyesight and vision. It is now known that beta-carotene is the primary precursor for Vitamin A in humans.

Moreover, the positive roles of beta carotene in maintaining a proper vision and its role in other degenerative diseases are now known [56]. There is plenty of evidence now that suggests carotenoids or food products rich in carotenoids are indeed beneficial in the prevention of several types of cancers and malignancies. Certain types of cancers namely, the cancers of prostate gland, urinary bladder and of the colon respond to a natural substance called lycopene. Carotenoid compounds has been used to treat various disorders such as erythropoietic protoporphyria. It is believed to act as an antioxidant and an immune system booster. Carotenoids contents tested in the bacterial extracts were Xanthophyll, β-cryptoxanthin and β-carotene (Table 5). Carotenoid compounds significantly varied among the two bacterial extracts; the highest total carotenoid concentration was shown in extract HA-MS-41 (1.165µg/ml). Whereas, lower values were measured in the HA-MS-118 [JX484801] extract (0.185 µg/ml) as shown in (Table 5).

**Table 5:** Concentrations of carotenoid compounds in the bacterial extracts HA-MS-41 and HA-MS-118.

**Flavonoid Compounds**

The presence of two flavonoid compounds was tested in the two bacterial extracts; Catechin and Rutin (Figure 6). Rutin compound was below the detectable limit in the tested two extracts. The highest catechin values were found in HA-MS-118 [JX484801] extract (8.652 mg/ml) as shown in (Table 6).

**Figure 5:** Structure of (A) Xanthophyll, (B) β-cryptoxanthin and (C) β-carotene.

**Figure 6:** Structure of (A) Catechin and (B) Rutin.

**Table 6:** Concentrations of flavonoid in the tested bacterial extracts.

| Extract no. | Concentration of catechin (mg/ml)* | Concentration of Rutin (mg/ml)* |
|-------------|-----------------------------------|---------------------------------|
| HA-MS-41 [JX484800] | 5.430 | N.D. |
| HA-MS-118 [JX484801] | 8.652 | N.D. |

*N.D.: not detected. All analyses were mean of triplicate measurements.*
Total Flavonoid Content: Flavonoids, among other biologically active compounds such as sterols, alkaloids, saponins, glycosides, and tannins support the significant bioactivity exhibited by our crude extracts including inhibitory properties against a wide range of pathogens. There is an increasing interest toward research on flavonoids occurring in various sources of marine organisms including sponges and their associated bacteria due to their multipurpose health benefits such as free-radical scavenging property, anticancer activity, antiviral properties etc [1-9,57]. Highly significant quantities of flavonoid contents were found in crude extract of the tested bacterial isolates equivalents of (+ catechin). The highest level of total flavonoid content was shown in HA-MS-41 extract. Whereas, the lowest value was shown in HA-MS-118 extract (Table 7).

Table 7: Total flavonoid contents in the bacterial crude extracts expressed as mg catechin equivalent / ml.

| Extract no.   | Concentration of flavonoid compounds (mg catechin equivalent / ml)* |
|---------------|-------------------------------------------------------------------|
| HA-MS-41 [JX484800] | 5.0±0.23                                                          |
| HA-MS-118 [JX484801] | 2.5±0.26                                                          |

*Results = Mean (n=3) ±S.E.

GC/MS Profiling for the Bacterial Crude Extract: The GC/MS analysis of crude bacterial extracts as shown in (Figures 7 & 8) indicates the presence of different bioactive components. The major components present in the ethyl acetate crude extracts are indicated in (Tables 8 & 9). Sponges are important source of functional components for the development of new chemotherapeutic agents. Phytochemical investigation of the ethyl acetate extracts of the studied sponge-associated bacteria revealed the presence of various phytochemical such as phenolic compounds, flavonoids, saponins, steroids, tannins. The bioactive composition of the two studied bacterial extracts obtained by GC/MS analysis and their bioactivity profiling are shown in (Table 10). Data revealed that the crude extract HA-MS-41 contained the maximum diversity of identified compounds (9 compounds). Whereas, HA-MS-118 extract contained the lowest number of the identified compounds (4 compounds). All the revealed compounds were identified and confirmed the presence of a variety of natural product families including mono- and di-terpenes, steroids, phenolics, flavonoids, carotenoids and fatty acids. The occurrence of different bioactive compounds suggests a wide range of biological application of the tested extracts.

Figure 7: GC-MS patterns of the bacterial ethyl acetate extract HA-MS-41 [JX484800].

Figure 8: GC-MS patterns of the bacterial ethyl acetate extract HA-MS-118 [JX484801].
Table 8: Bioactive compounds identified in the crude extract HA-MS-41 [JX484800] by GC-MS.

| Identified compounds* | RT (min) | Chromatogram % area | Chemical formula |
|-----------------------|---------|---------------------|------------------|
| Limonene              | 11.75   | 1.64                | \(C_{10}H_{16}\) |
| Bicyclo[2.2.1]heptan-2-one, 1,3,3-trimethyl- | 13.14   | 1.16                | \(C_{10}H_{16}O\) |
| Benzene, 1-methoxy-4-(1-propenyl)- | 16.16   | 0.03                | \(C_{10}H_{12}O\) |
| 12-Methyl-E,E-2,13-octadecadien-1-ol | 21.29   | 0.05                | \(C_{19}H_{36}O\) |
| 9,10-Secocholest-5,7,10(19)-triene, 1,3-diol, 25-[(trimethylsilyl)oxy]-, (3\(\Delta\)5Z,7E)- | 22.08   | 0.03                | \(C_{30}H_{52}O_3Si\) |
| \(\beta\)-D-Glucopyranoside, methyl 2-(acetylamino)-2-deoxy-3-O-(trimethylsilyl)-, cyclic methylboronate | 23.75   | 0.02                | \(C_{13}H_{26}BNO_6Si\) |
| Hexadecanoic acid, ethyl ester | 24.49   | 0.16                | \(C_{18}H_{36}O_2\) |
Table 9: Bioactive compounds identified in the crude extract HA-MS-118 [JX484801] by GC-MS.

| Identified compounds*                                                                 | RT (min) | Chromatogram % area | Chemical formula |
|-------------------------------------------------------------------------------------|----------|---------------------|------------------|
| Cyclohexanol, 1-methyl-4-(1-methylethenyl)-, acetate                                | 11.63    | 1.50                | C_{12}H_{20}O_{2} |
| Bicyclo[2.2.1]heptan-2-one, 1,3,3-trimethyl-                                         | 13.02    | 1.95                | C_{10}H_{16}O     |
| 3,5-Dimethyl-5-hexen-3-ol                                                            | 14.26    | 0.01                | C_{8}H_{16}O      |
| Benzene, 1-methoxy-4-(1-propenyl)-                                                   |          | 0.00                | C_{10}H_{12}O     |

*Number of identified compounds is (9).

Table 10: Bioactive composition of the ten studied bacterial extracts obtained by GC/MS analysis and their bioactivity profiling.

| No. | Compound                          | Chemical nature              | Extract No. | Bioactivity                                           | References |
|-----|-----------------------------------|------------------------------|-------------|------------------------------------------------------|------------|
| 1   | Limonene                          | Monocyclic-mono-terpene      | HA-MS-41    | +                                                    | [58-60]    |
|     |                                   |                              | HA-MS-118   | Anti-cancer antioxidant, growth supplement           |            |
| 2   | Bicyclo[2.2.1] heptan-2-one, 1,3,3-trimethyl- | Sesquiterpene Hydrocarbons | +                       | Anti-inflammatory, anticancer, anti-bacterial         | [61,62]    |
FT-IR Spectroscopy: FTIR spectroscopy was useful for the compound identification and when run under IR region in the range of 400-4000 cm\(^{-1}\) there was a variation in the peaks in the tested bacterial extracts. FTIR allows detecting the whole range of infrared spectrum simultaneously providing speed and accuracy in measurements of biological specimens. (Figures 9 & 10). Based on the functional group analysis, the bacterial extracts don’t contain any toxic compounds. Furthermore, the IR analysis gave results that suggest the presence of different functional groups ranging from O-H stretching, hydroxyl (3472.7-3362.6 cm\(^{-1}\)), C-H stretching, alkyl (2933.7-1460.6 cm\(^{-1}\)), C=C stretching (1667.3-1414.4 cm\(^{-1}\)), C-O bending, alcohols, ethers, esters, carboxylic acid and anhydrides (1382-1041.4 cm\(^{-1}\)), C=O stretching, carboxylic, carbonyl (1923.7-1721.3 cm\(^{-1}\)) and C-N bending, alkaloids (1405.2-1234.6 cm\(^{-1}\)).
Elemental Analysis: Elemental analysis of the tested extracts were presented in (Tables 11 & 12). The results of the elemental analysis of the ethyl acetate extract of the two tested bacteria indicated that the bacterial extracts contained major elements like Na, Mg, Al, Si, S, K, Ca, P and Cl. Besides, the major elemental constituents in all living organisms; C, N and O. The results showed considerable concentrations of many halides (Fe, Cu, Zn and Br). Extracts number HA-MS-118 showed the highest concentration percentages of Br (2.367 %), Fe (0.093 %), whereas extract HA-MS-41 showed the highest concentrations of Na (1.330 %) and K (2.297 %) (Table 11). Elements iron (Fe), copper (Cu), cobalt (Co), nickel (Ni), zinc (Zn), manganese (Mn), magnesium (Mg), molybdenum (Mo), chromium (Cr), vanadium (V), lithium (Li), selenium (Se), fluorine (F) and iodine (I) play both curative and preventive roles in fighting diseases of animals, plants and human beings [75]. The tested two bacterial extracts had high Ca and Mg contents. Calcium functions as a constituent of bones and teeth, regulation of nerve and muscle function. In blood coagulation, calcium activates the conversion of prothrombin to thrombin and also takes part in milk clotting. It plays a vital role in enzyme activation. It regulate heart rhythm, helps in normal blood clotting, maintain proper nerve and muscle functions and lower blood pressure.

Table 11: Elemental analysis of the ethyl acetate extract of the HA-MS-41 [JX484800] bacterial isolate.

| Element | Percentage (%) | Element | Percentage (%) | Element | Percentage (%) |
|---------|----------------|---------|----------------|---------|----------------|
| O       | 53.097         | Cl      | 0.907          | Na      | 1.330          |
| C       | 26.843         | S       | 0.857          | Al      | 0.063          |
| N       | 13.523         | P       | 1.143          |         |                |
| Ca      | 0.260          | Si      | 0.133          |         |                |
| K       | 1.467          | Mg      | 0.137          |         |                |

Table 12: Elemental analysis of the ethyl acetate extract of the HA-MS-118 [JX484801] bacterial isolate.

| Element | Percentage (%) | Element | Percentage (%) | Element | Percentage (%) |
|---------|----------------|---------|----------------|---------|----------------|
| O       | 41.373         | Ca      | 0.117          | Mg      | 0.100          |
| C       | 34.627         | K       | 2.297          | Na      | 1.053          |
| N       | 17.173         | Cl      | 0.270          |         |                |
| Br      | 2.367          | S       | 0.370          |         |                |
| Fe      | 0.093          | P       | 0.230          |         |                |

Mg is also an essential activator for the phosphate-transferring enzymes myokinase, diphosphopyridine nucleotide kinase, and creatine kinase. It also activates pyruvic acid carboxylase, pyruvic acid oxidase, and the condensing enzyme for the reactions in the citric acid cycle. It is also a constituent of bones, teeth, enzyme cofactor; kinases, etc. [76]. In the current study, among the various minerals potassium was observed also in high level in both tested two extracts. Without potassium which is an essential for the transport for nutrients inside the cell, nutrients could not be able to enter into the cell that leads cell death. Potassium is followed by phosphorous in the tested two extracts. Sodium, on the other hand, is the principle extracellular cation used for osmoregulation in inter modular fluid of human body was measured in the two tested extracts. The highest Na content was obtained from the HA-MS-41 bacterial isolate.

Silicon which is also another important element to prevent the hardening blood veins and arteries was observed in the HA-MS-41 bacterial isolate, but not in the HA-MS-118 bacterial isolate. Al was measured in the HA-MS-41 bacterial isolate, but not in the HA-
MS-118 bacterial isolate. In the meantime, Br and Fe were found in the HA-MS-118 bacterial isolate, but not in the HA-MS-41 bacterial isolate (Tables 11 & 12). Iron is an essential element because it is important in several enzyme reactions as co-factors. It has an oxidizing effect, it acts as astringent. The current study indicated the presence of considerable amounts of halogens such as chlorine and bromine. The presence of halogen substituents in many natural products profoundly influences their biological activity. Chemical structures of marine products often differ from terrestrial secondary metabolites in being halogenated with bromine and/or chlorine. Chlorine is found in the HA-MS-118 bacterial extract.

Unlike terrestrial organisms, marine organisms often produce halogenated secondary metabolites, particularly alkaloids [77]. The results of the current study showed that the highest bacterial extracts in bromine content was HA-MS-118. It has been suggested that marine life produces most of the 4,000 known natural organohalogens, and almost all of the 2,100 natural organobromine compounds are found in marine organisms [78]. Halogen containing secondary metabolites exhibit interesting biological effects, such as antibiotic, anticancer and others. On the other hand, (Tables 11 & 12) showed that the two tested bacterial extracts contained sulfur in variable amounts, among the highest bacterial extracts in sulfur contents was the HA-MS-41 bacterial isolate. Sulfur in the form of organic and inorganic compounds is always present in all living organisms and is an important biogenic element. The average content of sulfur is relatively higher in marine organisms. On average natural product contain less nitrogen, sulfur, and halogen atoms but in marine natural product there is higher ratio of this constituent. Moreover, sulfur containing phenolic compounds can be reasonably speculated as strong anti-inflammatory and antioxidant agents. Activity of sulfur-containing phenolic antioxidants in vivo is mediated by their effect on redox-sensitive transcription factors [79]. Conclusively, the presence of these elements in the tested extracts marks its usage in therapeutic purposes and may remark the tested bacterial extracts in the current study as promising bioactive materials.

**Biochemical Assays**

**Antioxidant Capacity:** Living cells in the human body continuously produce reactive oxygen species and free radicals and are controlled by body enzymes. Over production of either free radicals, reactive oxygen species or failure in the defense mechanisms result in severe damage to cells and tissues leading to non-communicable chronic diseases, specifically cancer [80]. Antioxidants are capable of inhibiting the oxidation that produces free radicals. In turn, these radicals can start chain reactions that damage cells. Antioxidants terminate these chain reactions by removing free radicals intermediates and inhibit other oxidation reactions [81]. In the current study, the levels of total antioxidant capacity (TAO) were measured in the bacterial extracts using DPPH, ABTS, NO and Phosphomolybdenum assays. The present results revealed a potential antioxidant activity of the two tested bacterial extracts.

**DPPH Scavenging Activity:** The results of DPPH test indicated that IC50 values of ethyl acetate extracts HA-MS-41 and HA-MS-118 bacterial isolate were lower than the standard used for the assay; ascorbic acid (IC50 = 2.52 mg/ml). DPPH scavenging activity in terms of IC50 values was shown to be high in case of extract HA-MS-41 (1.045 mg/ml) and in case of extract HA-MS-118 (1.473mg/ml). This may be due to the presence of some potent antioxidant compounds such as limonene, amiodarone and 3',8,8'-Trimethoxy-3-piperidyl-1,2',2''-binaphthalene-1',1''4,4'-'tetrone which are powerful antioxidants and have good radical scavenging activities. This finding is also compatible with the total phenolic and flavonoids contents of the two tested extracts (Table 13).

**Table 13:** Total antioxidant of the examined two extracts using DPPH assay.

| Extract concentration (mg/ml) | TEAC/mM equivalent Ascorbic acid* |
|-----------------------------|----------------------------------|
|                             | HA-MS-41 [X484800] | HA-MS-118 [X484801] |
| 30                          | 94.69±0.01           | 89.74±0.04          |
| 6                           | 93.13±0.001          | 91.03±0.005         |
| 1.2                         | 93.95±0.03           | 85.93±0.07          |
| 0.24                        | 88.88±0.12           | 66.49±0.03          |
| 0.048                       | 50.53±0.49           | 29.69±0.022         |
| 0.01                        | 48.71±0.78           | 9.79±0.11           |
| IC50 (mg/ml)                | 1.045               | 1.473               |

*Data are mean ± SD

**Nitric Oxide (NO) Assay:** Interestingly, the antioxidant potency of the tested bacterial extracts was also measured by NO assay; since NO- has a damaging effect as RNS in in vivo system. Nitric oxide (NO-) is radical compound produced during the transformation of L- arginine to citrulline by nitric oxide synthase (NOS). Nitric oxide functions as the effecter molecule for macrophage-mediated cytotoxicity and has also been recognized to inhibit mitochondrial respiration and DNA synthesis. Nitric oxide recently raised a lot of interests as a suppressive factor that inhibits anti-tumor immune response. Readings are based on the colorimetric change, which occurs when naphthyl ethylenediamine is added to the by-product of reaction between sulfanilamide and nitrite. Highly significant TEAC value was obtained (Table 14). Percentage inhibition gradually increased with increase in concentration. IC50 values of

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the antioxidant activity of bacterial extracts using NO assay for the tested two extracts were 5.98 and 6.22 %. That was in agreement with the high phenolic and flavonoid compounds present in the tested extracts [81].

**Table 14**: Total antioxidant of the examined two extracts using Nitric oxide (NO) assay.

| Extract concentration (mg/mL) | TEAC/mM equivalent Ascorbic acid* |
|-------------------------------|---------------------------------|
| HA-MS-41 [JX484800]          | HA-MS-118 [JX484801]            |
| 30                            | 72.15±0.013                     | 70.47±0.006                     |
| 6                             | 59.91±0.005                     | 37.48±0.111                     |
| 1.2                           | 31.19±0.004                     | 39.81±0.01                      |
| 0.24                          | 23.36±0.003                     | 33.27±0.004                     |
| 0.048                         | 21.4±0.013                      | 32.52±0.001                     |
| 0.01                          | 18.66±0.027                     | 31.5±0.002                      |
| IC₅₀ (mg/ml)                  | 5.98±0.003                      | 6.22±0.012                      |

*Data are mean ± SD

**ABTS+ assay**: DPPH assay was confirmed by ABTS assay. Highly significant TEAC values were obtained in the tested bacterial extracts (Table 15). Percentage inhibition gradually increased with increase in concentration. The results of the current study cleared out that the ABTS assay gave completely different results for the antioxidant activity measured in the tested two extracts (Table 15). The percentage inhibition of the HA-MS-41 [JX484800] was higher (IC₅₀ = 3.40 mg/ml) than that of the HA-MS-118 [JX484801] extract (IC₅₀ = 1.64 mg/ml) revealing much higher antioxidant capacity. The high potency of the two extracts as strong antioxidant capacity may returns to its high phenolic contents (4.991 & 10.369 µg/ml, respectively), specially caffeic acid (1.165 5.642 µg/ml, respectively).

**Table 15**: Total equivalent antioxidant activity and IC₅₀ values of the bacterial extracts using ABTS+ assay.

| Extract no. | Conc. (mg/ml) | % inhibition | mM Trolox equivalent | IC₅₀ (mg/ml) |
|-------------|---------------|--------------|----------------------|-------------|
| HA-MS-41 [JX484800] | 30             | 94.34        | 8.53                 | 3.40        |
|             | 6              | 68.77        | 5.67                 |
|             | 1.2            | 39.8         | 2.43                 |
|             | 0.24           | 25.72        | 0.86                 |
|             | 0.048          | 22.16        | 0.46                 |
|             | 0.01           | 20.54        | 0.28                 |
| HA-MS-118 [JX484801] | 30             | 88.51        | 7.88                 | 1.64        |
|             | 6              | 66.02        | 5.36                 |
|             | 1.2            | 63.59        | 5.09                 |
|             | 0.24           | 52.10        | 3.81                 |
|             | 0.048          | 37.80        | 2.22                 |
|             | 0.01           | 26.37        | 0.93                 |

**Phosphomolybdenum assay**: Total antioxidant capacity of the ethyl acetate extracts using the Phosphomolybdenum expressed as the number of equivalents of ascorbic acid and based on the reduction of Mo (VI) to Mo (V) by the antioxidant compound and the formation of a green phosphate/Mo (V) complex with a maximal absorption at 695 nm revealed that the antioxidant activity of the tested two extracts exhibit increasing trend with the increasing concentration of the bacterial extract (Table 16). Addition of various concentrations of the extracts showed that extract HA-MS-118 [JX484801] (37.75±9.23 mM equivalent ascorbic acid at extract concentration 30 mg/ml) was more effective to reduce Mo (VI) to Mo (V), while the lowest effects were shown by HA-MS-41 [JX484800] (25.53±0.74 mM equivalent ascorbic acid at extract concentration 30 mg/ml) (Table 16). The reduction of Mo (VI) to Mo (V) suggested the presence of effective antioxidants in HA-MS-118 [JX484801] and HA-MS-41 [JX484800] bacterial extracts (Table 16).

**Table 16**: Total antioxidant of the examined two extracts using phosphomolybdenum assay.

| Extract concentration (mg/ ml) | TEAC/mM equivalent Ascorbic acid* |
|-------------------------------|---------------------------------|
| HA-MS-41 [JX484800]           | HA-MS-118 [JX484801]            |
| 30                            | 25.53±0.74                      | 37.75±9.23                     |
| 6                             | 6.43±0.097                      | 7.16±3.76                      |
| 2.5                           | 2.65±0.035                      | 1.74±0.13                      |
Anticancer Assays

Cytotoxicity Studies: For many years past, the cytotoxic actions of natural products were regarded as the first step in identifying cancer chemotherapeutic drugs. In vitro cytotoxicity assays are a potentially useful tool in the study of toxic compounds of complex mixtures. A significant number of marine compounds have shown potential anticancer activities against the different hallmarks of cancer, such as cell growth inhibition, antimitotic activity (anti-tubulin effects), apoptosis and/or autophagy induction, and migration, invasion or metastasis inhibition. For most of the isolated compounds and their semisynthetic derivatives, the reported activities have been observed in vitro, but a significant number of these compounds, i.e., more than twenty, are either approved or in clinical phase. MNPs exhibit extensive chemical variability and complexity and include alkaloids, polyketides, terpenes, peptides and carbohydrates. Although marine resources are also limited, sourcing issues should be solved in the future by using aquaculture and biotechnology processes and the improvement of chemical synthesis in order to provide sufficient amounts of compounds for research purposes or pharmaceutical development [82].

In the present study, the tested crude extracts of the bacterial isolates, each at six different concentrations (50 – 1.56 μg/ml), were used to screen the cytotoxicity against breast cancer (MCF7), colon cancer (HCT) and hepatocellular carcinoma (HepG2). It has been observed that the two extracts obtained from the tested sponge associated bacteria were toxic against those carcinoma cells (Table 17). The present results revealed that the maximum cell growth inhibitory effects of bacterial crude extracts against the tested cancer cell lines; HepG2; HCT and MCF-7 were obtained from extract HA-MS-118 (IC50= 36.5; 23.6; 11.9 μg/ml, respectively). While, the minimum IC50 values against all the tested cancer cell lines HepG2; HCT and MCF-7 were obtained from the extract HA-MS-41 (IC50= 22.7; 14.6; 12.2 μg/ml, respectively (Figures 11 &12) show the cytotoxicity trends of the bacterial extracts against the three cancer cell lines. These values represent in all cases a comparable high toxicity to many other marine bacteria crude extracts Figures The results of this study are in good agreement with previous findings reported by Abdelnasser et al. [83]. Interestingly, the comparatively high cytotoxic effect of the extract HA-MS-118 may be due to its high Boron and Fe contents along with the presence of reasonable amounts of different biologically active compounds. Moreover, data obtained from cytotoxicity test against cancer cell lines are compatible with the high amounts of phenolic, flavonoids, alkaloids and other anticancer natural compounds shown in the tested extracts.

Table 17: IC50 values of the bacterial extracts against the three tested cancer cell lines.

| Cancer cell line | Bacterial extract | IC50 of the bacterial crude extracts (μg/ml) |
|------------------|-------------------|---------------------------------------------|
|                  | HA-MS-41          | HA-MS-118                                   |
| HepG2            | 22.7              | 36.5                                        |
| HCT              | 14.6              | 23.6                                        |
| MCF-7            | 12.2              | 11.9                                        |

Figure 11: Cytotoxic activity of extract HA-MS-41 against cancer cell lines; (A): MCF-7, (B): HepG and (C): HCT.
Figure 12: Cytotoxic activity of extract HA-MS-118 against cancer cell lines; (A): MCF-7, (B): HepG and (C): HCT.

Protein kinase Inhibitory Effect

Previous cytotoxicity data were confirmed by a specialized screening of anticancer test; protein kinase inhibition effect. Protein tyrosine kinases (PTKs) are key enzymes in many signal transduction pathways and play a crucial role in many cell regulatory processes. Functional perturbation of some of the enzymes results in a variety of diseases including cancer and inflammation [84,85]. This study (Table 18) revealed that the highest PTK inhibition activity was obtained from extract HA-MS 118 (inhibition % = 94.24±1.15). Whereas, the lowest inhibition activity was obtained from extract HA-MS 41 (inhibition % = 87.93±0.51) indicating high potency of inhibition activity. This may be due to the presence of natural product compounds such as flavonoids and alkaloids which are considered as protein kinase inhibitors [86]. It has been stated that marine sponge metabolites had proven to be extremely potent against a range of kinase targets heavily involved in an increasing list of disease mechanisms including cancer, Alzheimer’s disease and atherosclerosis [82,87,88].

Table 18: Effect of the HA-MS-41 and HA-MS-118 bacterial extracts on protein kinase inhibition.

| Extract  | Inhibition % * |
|----------|----------------|
| HA-MS-41 | 87.93±0.51     |
| HA-MS-118| 94.24±1.15     |

*Data are means ± S.D.

Conclusion

In continued search in our laboratory for novel microbial secondary metabolites, bacterial strains were screened. This has resulted in isolation of a promising strain of Proteobacteria species from the Red Sea sponge *Ircinia echinata*. Two tested isolates, HA-MS-41 and HA-MS-118 showed antioxidant activity as well as cytotoxicity against three established cancer cell lines. This study cleared out the importance of bioprospecting studies of marine biodiversity for bioactive natural compounds discovery and development of new drugs. The active extracts deserve special attention in further studies on the isolation and structure determination, as well as more refined biological assays; since clearly, the bacteria associated Red Sea sponge assayed could play an important source for the development of new biotechnological products.

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