EXTRACTION, ISOLATION OF ACTIVE PRINCIPLES, ANTI-BACTERIAL AND WOUND HEALING ACTIVITY OF THE MARINE ALGAL SPECIES OEDOGONIUM GLOBOSUM AND OEDOGONIUM INTERMEDIUM

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ABSTRACT

Objective: Algae is the undisputed treasures of the sea and are a valuable raw material, providing unlimited opportunities for new drug discoveries. Marine algal products are in demand in the international market in the form of standardized algal extracts or semi-finished products.

Methods: Aqueous and methanolic extracts of Oedogonium globosum and Oedogonium intermedium species were obtained maceration and hot percolation. The active principles from O. intermedium were isolated, purified by column chromatography, and characterized by spectral studies IR, λmax, 1HNMR and MS. The extracts of Oedogonium species were screened for their anti-microbial effects, acute dermal irritation and wound-healing activity studies.

Results: Comparing to Oedogonium intermedium (45%, 90%, 87%), very low extractive yields were obtained for Oedogonium globosum (10.80%, 37%, 28%). At phytochemical screening, Terpenoids, Flavonoids and Glycans were found to be present in a significant amount and upon their isolation, it was found that a collection of fractions from cold extract with RI value in the range 0.32-0.34 as Flavonoids and those from the hot extract with 0.40-0.72 as Flavonoids and those from methanolic extracts with 0.23, 0.44 and 0.71 as for Terpenoids. Anti-bacterial studies revealed out the fact of Oedogonium species could give higher inhibition to gram-positive than for gram-negative bacteria at (10 μg/10μl/disc) concentration. No symptoms of systemic toxicity and mortality were observed. Silver sulfadiazine, more potent in wound closure, the effect of methanolic extracts of O. intermedium (87%) was almost at par to the standard (95%) in action and significantly greater than O. globosum (72%, P<0.05).

Conclusion: Admittedly, Oedogonium type algal species can be known as medicinal algae with a plethora of a wide range of pharmacological activities. Thus, this research work may be considered further for extensive innovative discoveries of new lead molecules and any other pharmacological activities, in the future.

Keywords: Oedogonium Globosum, Oedogonium Intermedium, Extraction, Phytochemical analysis, Isolation, Dermal irritation test, Anti-microbial and Wound Healing Activities

INTRODUCTION

Nowadays, esteemed researches involve the transformation of these sponges into rich value biomass co-products in the pharma, nutrition, topical, cosmetic and greenery fields, livestocks, and organic fertilizers also, 50% of all oxygen production is by these nutrition, topical, cosmetic and greenery fields, livestocks, and organic fertilizers also, 50% of all oxygen production is by these

From the literature, the marine sponges do contain many chemicals viz, terpenes, sesquiterpenes, terpenoids and steroids, polysaccharides, essential amino acid, vitamins, minerals, cyclic-and peptides, proteins, alcohols, tannis and phlorotannins, lipids, fatty acids, nucleic acid bases, ribonucleic acid (RNA), de-oxiribonucleic acid (DNA), macrolides, polyketides [3-5]. It’s a good source of minerals, vitamins, proteins, carbohydrates, and fibers [6, 7]. The proficient biological activities of algae from ancient times to a modern era include antidiabetic, anticancer [8, 9], antivirals, antimicrobials, anti-inflammatory [10], antioxidant [11], anticoagulants, antibiotics [12], antihypertensive agents, blood cholesterol reducers, dilutory agents [13], insecticides and antithromogenic agents [14], nutritional supplement, for increasing skin thickness and elasticity beauty enhancers, growth factor modulators and other industrial purposes.

Since Phyto-constituents are multicomponent mixtures, their purification and determination are still very difficult. Wound closure property is incomparable in herbals and marine seaweeds and that may be by the enhanced formation of granulated tissues, collagen, skin appendages, and re-epithelialization [17]. On all the above background literature studies and analyses, in this paper, we are more interested to report first about its extraction results, microbialid and wound ameliorating effects of the methanolic and aqueous extracts of Oedogonium globosum and Oedogonium intermedium and secondly on the isolation and characterization of active principles from Oedogonium intermedium.

MATERIALS AND METHODS

The chemicals and reagents used for the study, were of analytical grade and were procured from E-Merck. In this research project, some of the instruments viz., Schimadzu UV spectrophotometer (Model No. UV-2400PC) for UV spectral study, PC-FTIR equipment (Make: Schimadzu 8201) for IR spectral study and a Bruker 400 machine for 1HNMR and 13CNMR spectra were made used of.

Collection and authentication of plant material

Two freshwater seaweed samples were collected from local and in and around the regions of our institution, i.e. Othakalmandapam, Malumichampatti and Kinathukkadavu, Coimbatore, Tamil Nadu, during the low tidal conditions at a depth of 1.3 m. Unsuitable matter viz, epiphytes, extraneous matter, and necrotic parts of the collected samples were removed. Algal samples with enough
quantity were cleaned subsequently with tap-, fresh-and sterile distilled water, let the shade dried, chopped and crushed into a powder form in a mortar and pestle. Herbarium sheet was prepared and the taxonomic position of the same was identified by BSI, Southern Regional Centre, TNAU Campus, Coimbatore (BSI/SRC/05/23/2017/Tech./3620 dated 29.03.2017). This report stated the scientific classification of the selected algae as Plantae kingdom; Chlorophyta division; Chlorophyceae class; Oedogoniales order; Oedogoniaceae family; Oedogonium genus and Globosum and Intermedium species.

Extractions
Algal extracts were prepared by maceration and hot extraction method. After drying the leaves of both algae (1 Kg each) in the working lab for 1 w, they were milled and ground into a fine or very nice coarse powder (100 g) using a miller. The dried material of both Oedogonium globosum and Oedogonium intermedium was weighed and the powdered algal sample (100 g) was put into the extractor (1000 ml capacity) with solvents (500 ml each) of increasing polarity viz., chloroform, methanol, and cold water, hot water for a duration of 2 h, 3 h and 4 h for non-polar, cold and hot-polar solvents respectively. The contents were shaken at regular intervals and the solvent extract thus obtained was concentrated at 40 °C to dryness to get a greenish crude product. Extractive values and the yield percentage for all categories were estimated and compared.

Phytochemical screening
Secondary metabolites present in the plants/algal extracts can be identified by a fast, simple, and inexpensive procedure of Phytochemical screening, which can be performed with an active fraction of algae by using the appropriate standard tests [18].

Phytochemical isolation
From the literature reports [19], it was confirmed that not much research work has been done regarding the bioactive compounds from Oedogonium intermedium. All the three solvent extracts viz., cold- and hot-water extracts and methanol extracts were made dry by suction and kept in a hot air oven set at 50 °C. They were then mixed with pre-warmed silica and introduced into three separate columns (first one packed with methanolic extract, a second column with hot aqueous and the third one with cold aqueous extract of Oedogonium intermedium), each charged with silica gel, 60-120 mesh (500 g) for chromatographic separation. The columns were eluted with different solvents in order of increasing polarity. The elutes, in serial, had been subjected for the thin-layer chromatographic technique for the identification of the isolated samples through Rf value estimation. The fractions, having similar Rf values, were merged together, evaporated to dryness using rotavapor, and recrystallized with respective eluting solvents of methanol and distilled water.

Anti-microbial study
Minimum inhibitory concentration study by disc diffusion mechanism
The sterile impregnated disc, with diluted aqueous and methanolic extracts of algal species at varying quantities (100, 50 and 25 µg/ml), was placed on the agar surface. A loop full of the organisms at 10⁶ cfu/ml quantity inoculated the incubated plates at 37 °C. The size (diameter) of the inhibition zones was observed and calculated [20].

Zone of inhibition study
The suspension of the test microorganisms 10 µl (10⁶ cells/ml) was applied using a sterilized cotton swab by spread plate method, having the bacterial cultures in Muller Hinton agar media. After solidification, the filter paper discs were impregnated with the test samples of the extracts and placed by sterilizing applicators on test organism plates. Compounds were screened for anti-bacterial activity against Escherichia coli (ATCC. No. 25922) and Staphylococcus aureus (ATCC. No. 51740) using Amikacin (100 µg/ml) as standard and distilled water (100 µg/ml) as control. After incubating the assay plates at 37 °C for 24 h, calculated the width of growth inhibition zones. The determination was performed in triplicate [21].

Wound-healing pharmacological activity
The wound-healing activity of aqueous extracts of Oedogonium globosum and Oedogonium intermedium was studied by excision wound model, following the international ethical guidelines and under the supervision of the scrutinizing committee of Institutional Animal Ethical Committee (IAEC), Karpagam University (ethical approval No.106, Oct 10, 2011).

Animal selection and acclimatization
Institutional animal ethical committee approval, reference No. KU/IAEC/Ph. D/065 has been obtained for our in vivo experiments. Both male and female Wistar Albino rats (150-200 gm) were purchased from Amrita Institute, Kerala and were maintained in the following manner, in our college animal house for 7 d, housed in polypropylene cages, fed with quality rodent pellet diet, water ad libitum, room temperature 21-25 °C, relative humidity 50-60 % and kept for fasting over night for at least 12 h. Feeding tubes and syringes were used to administer the drug.

Acute dermal irritation test, OECD guidelines 404
Samples (2 mg/kg) were prepared in ointment form with olive oil, for dermal application. A standard irritant patch (0.8% w/v formaldehyde), a control (placebo) patch and test samples of aqueous extracts of Oedogonium globosum-and Oedogonium intermedium-loaded transdermal patches were prepared for the study. An area of approximately 150 cm²(10x15 cm²) was exposed to aid the scoring by removing the dorsal fur [22].

Test sample (0.5 ml) was applied to a metallic patch with one rat, mounted on a Micropore tape, wrapped around the abdomen and secured by an elastic bandage as an initiation of the study. When once the dressing was removed, no sign of necrosis or corrosion was observed. The same procedure was continued with two samples to separate skin-sites and or with two further animals. Dermal response scores at 1, 24, 42 and 72 h were evaluated for each animal, then after for the next 7 d as a further observation. Untreated skin of adjacent areas was treated as a control. The irritation scores of erythema and edema were scored on a scale of 0-4 and to obtain the primary irritation index; mean scores were averaged.

Excision wound model
Animals were anesthetized at the time of wound creation. An excision wound of circular area 500 mm² and 0.2 cm depth, marked...
The animals were divided into four groups (n=6). The group I animals (placebo control) were topically treated with Ointment base I. P., Group II and Group III with the 10% ointment of the aqueous extract of Oedogonium globosum and Oedogonium intermedium respectively; and Group IV with the 10% ointment of the standard, Silver sulfadiazine. The treatment procedure, dressing up the wounds with standard and test samples every day, was started immediately after the wound creation for 14 d. The wound closure rate was assessed using a graph paper on days 0, 3, 5, 7, and 14 post-wounding. The period of epithelialization was done by measuring the % of reduction in the wound area size at intervals from the next day of treatment.

Wound healing % = \left( \frac{1 \text{ st day wound spot area} - \text{Specific day wound spot area}}{1 \text{ st day wound spot area}} \right) \times 100

The data were applied to one-way ANOVA and Dunnett’s multiple comparisons to study the changes; P values significant criteria; Mean of result data by SEM Software: GraphPad Prism 5.01.

RESULTS

Two species of algae were extracted by cold maceration and hot extraction with distilled water, methanol and, chloroform. Since O. intermedium samples were found to be yielding more quantity of extracts (65 %) comparing to O. globosum (30 %, owing to dampened nature), we started to continue our researc h with the former algal species of the three extractions of Oedogonium intermedium, the yield of cold methanolic extract (72 %) was found to be significantly greater than hot methanolic (33 %). Similarly, comparing to the hot aqueous extract (61 %), the cold extract (70 %) was found to be more in yield (table 1). On phytochemical screening, glycans, flavonoids and terpenoids were to be present (table 2). TLC was performed on the extracted materials of algal species.

Table 1: Percentage yields of crude extracts

| S. No. | Name of the sample | Quantity of powdered material (g) | Name of the extract | Amount of extract (g) | % Yield | Melting point in °C |
|--------|-------------------|----------------------------------|---------------------|----------------------|---------|--------------------|
| Trial extraction | | | | | | |
| 1 | Sample-A (Oedogonium globosum) | 5 | Chloroform | 0.54 | 10.80 | - |
| 2 | Sample-B (Oedooonium intermedium) | 5 | Distilled water | 1.42 | 28.40 | - |
| 3 | Sample-B | 25 | Hot Methanol | 11.25 | 45.00 | - |
| 4 | Sample-B | 25 | Methanol | 22.50 | 90.00 | 210 |
| 5 | Sample-B | 25 | Hot water | 15.425 | 61.70 | 116 |
| 6 | Sample-B | 25 | Cold water | 21.75 | 87.00 | 67 |

Table 2: Preliminary phytochemical analysis of the aqueous and methanolic extracts of Oedogonium Intermedium

| S. No. | Phytoconstituents | AEOI | MEIO |
|--------|-----------------|------|------|
| 01. | Alkaloids | + | - |
| 02. | Polysaccharides | +++ | +++ |
| 03. | Saponins | - | - |
| 04. | Terpenoids | +++ | +++ |
| 05. | Flavonoids | +++ | ++ |
| 06. | Steroids | + | - |
| 07. | Glycosides | ++ | ++ |
| 08. | Tannins | + | - |

Where AEOI-Aqueous extract and MEIO-Methanolic extracts of Oedogonium intermedium; +++-highly present; ++-moderately present; +-slightly present and - absence of Phyto-constituents.

The various extracts were then separated for individual Phyto-constituents by the varying proportion of methanol and distilled water. When detected in iodine chamber and UV cabinet, single spotted elutes were collected and combined. Three different fractions were in such a manner obtained as pure Phyto -constituents (table 3).
Phyto-constituent I, Glycans: was obtained with cold distilled water as the eluent. A variety of glucoside derivatives of both (+) and (-) enantiomers in the form of pure green colored and amorphous powder have been isolated upon re-crystallization. Its TLC study revealed its presence through a single spot (Rf value 0.33) in the mobile phase of methanol and distilled water (3:7). A Melting point value is 60° C. IR spectra were characterized as IR: 3500-2500 cm⁻¹ for OH stretching; 1600 cm⁻¹ for C=C stretching, benzene nucleus; 1320-1210 cm⁻¹ C-O stretching, Aryl; 585 cm⁻¹ C-H bend, medium; The absorption maxima, λmax was recorded as 291 nm with distilled water as solvent. They were analyzed in the 1HNMR spectrum (δppm, D₂O) presence of four singlets at δ 1.182 (1H, CH₃,S), 1.224 (1H, CH₃,S,J=4.6Hz), 1.240 (1H, CH₃,d) and 1.813 (1H, CH₃,S) for three protons each were assigned to methyl groups. The sharp singlet observed at δ 1.183 (1H, CH₃,S) and 8.352 (1H, COOH,S) were assigned to –CH₃ and–COOH groups, respectively. Oxygenated proton afforded a doublet and a double triplet at δ 3.106 (1H, OH, d,J=11.5Hz) and 3.22 (1H, OH, dt). Other protons were as multiplet, lie in the range of 1.21-2.20. Mass (m/z): Base peak value is 90.87, the molecular ion peak has m/z value as 68.94 and the other peaks correspond to ion fragments, whose m/z ratios were found to be as 127.77, 154.66 and 184.69.

Phyto-constituent II, Flavanoids: was obtained with hot distilled water as the eluent. A pure amorphous powder of green coloured isolate had exited upon re-crystallization. Rf value is 67° C. It had been isolated upon re-crystallization. Its TLC study revealed its presence through a single spot (Rf value 0.77) in cyclohexane: acetone (–) enantiomers in the form of pure green colored and amorphous powder have been isolated upon re-crystallization. Rf value is 67° C. The absorption maxima, λmax was recorded as 290 nm with distilled water as the eluent. A pure amorphous powder of green coloured isolate had exited upon re-crystallization. Rf value is 67° C.

Well-defined erythema and very slight oedema have resulted in the treated skin-areas of the three rats, when exposed to 0.5 ml of test solutions (Oedogonium globosum and Oedogonium intermedium) for four hours. The skin irritation had resolved within 7days and no evidence of either a corrosive effect or systemic toxicity symptoms and or any mortality (table 5). Anti-oxidant activity, anti-cancer and anti-diabetic activities may be expected owing to polyphenolic compounds and astringent and anti-microbial properties owing to flavonoids. The current research’s results showed good wound closing responses which might be due to enhanced formation of granulated tissues, collagen, skin appendages and re-epithelialization. The algal extracts should have stimulated the endogenous antioxidants and thus protected the tissues from free radical-mediated damages.

Excision wound healing by contraction and epitelialization model was followed to assess the percentage of wound closure or closure rate study. Wound healing study data of the sponges has been shown in (table 6). Silver sulfadiazine showed more potent significant wound breaking strength compare to that of control group rats. Comparing to the control, all test samples showed a diminish in the area of the cut wounds on the 14th day (P<0.05). On the third and sixth days, the wound healing effect of the standard (46 % and 65 % respectively) was significant (P<0.05). The rate of wound closure of the test samples O. globosum (42 %) and O. intermedium (47 %) were found to be almost equal to that of the standard from 9th day of observation.
the gram-positive bacteria because of the complex structure of gram-
these algal species were found to be significant in action only against
and inhibitory action of the tested species, but was observed that Karpagam College of Pharmacy, Coimbatore–
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facilities to accomplish this research work successfully.

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acid, phosphotidyl serine isomeric derivatives and sulfoquinovosyl

halogenated hydrocarbons [31-32].

compounds like terpenes, acetogenins, indoles, phenols, fatty acids,
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The prevalence of anti-microbial chemical defence effects, effectively
observed in the seaweeds [30], may be due to the presence of chemical
compounds like terpenes, acetogenins, indoles, phenols, fatty acids,
halogenated hydrocarbons [31-32].

The inherent wound healing property of algae were going high in
value of wound healing property in all 14 d observation and the effect of O. intermedium [87 %] at par to the standard [95 %]
in action and significantly greater than the other O. globosum [72 %, P<0.05]. From literature, it was observed that the wound healing activity of seaweeds might be due to the proliferation of cells, collagen, granulation tissue formation [33]. Hydroxylated cinnamic acid, phosphotidyl serine isomeric derivatives and sulfouquinovosyl
diaclyglycerol [34-35] are the chemical constituents being found in
algal species, might be responsible for the proliferation and migration of keratinocytes and fibroblasts. These findings thus help us, the researcher, to proceed further with the later species for other related pharmacological evaluation like anti-coagulant or anti-
cancer activities in the future.

CONCLUSION

Thus, this current work adds to the reliability of this natural molecule
as a good candidate for medical use in the future in Indian folk
medicine. Until recently, there was a lack of reports on phytochemical
composition and other parts of either pharmacological actions or
standardization data or Physico-chemical properties details. Though
the algal species are, in general, quite promising for further work in
this regard, yet research can be keenly focused in the following aspects: i) In addition to extraction, the structure of the identified bioactive compounds, in the future, to be elucidated using other
spectroscopic determinations ii) Vast experimental data should be
retrieved from traditional medicines. iii) Modern scientific validation
and standardization methods can be encouraged to obtain a significant
market value of these marine species.

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DISCUSSION

Worldwide diseases’ responsible, foodborne pathogens, though show
resistance against contemporary antibiotics, found to be controlled by
more eco-friendly, natural products, being found in consumers’ diets
i.e. seaweeds [26]. Two species of algae were extracted by cold
maceration and hot extraction with distilled water, methanol and,
chloroform. In anti-bacterial study, O. intermedium was showing a comparatively higher inhibition effect than O. globosum. However,
these algal species were found to be significant in action only against
the gram-positive bacteria because of the complex structure of gram-
negative bacteria [27-29]. Though the data supports for the less
inhibitory action of the tested species, but was observed that E. Coli
and S. aureus organisms responded to the actions of algal isolations.

All values are by Mean ± SD (n=3). ***P<0.001, **P<0.01 and *P<0.05, as compared to control. One-way ANOVA followed by Turkey’s post hoc
multiple comparison tests.

AUTHORS CONTRIBUTIONS

M. Karpakavalli originated the research work and wrote the manuscript; A.Y. Sanglimuthu made suggestive revision; G. Prakash,
P. Sivasubramaniam and D. Renjithkumar developed the document;
finally S. Mohan has corrected the manuscript and approved the final
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CONFLICTING OF INTERESTS

Nil

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