In vivo Evaluation of Antimicrobial Effect of Methanolic Extract of Chlorella vulgaris on Impetigo and Some Dermatophytes

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**IMPETIGO** is one of the infectious superficial bacterial diseases and its treatment with antimicrobial agents may cause serious problems. So, a test of new microbial infection-fighting natural compounds is necessary. In the present study, histological examinations of the experimental animals revealed the effectiveness and safety of *Chlorella vulgaris* methanolic extract ointment used in the treatment of impetigo, tinea corporis and cutaneous candidiasis without any side effects on skin tissues. In addition, examination of skin sections treated with *C. Chlorella vulgaris* extract possessed no significant toxic effects. The skin appeared with normal epidermis as the keratinized fibers of stratum corneum were regularly arranged, appeared condensed without any disruption. The dermis appeared normal with minimal inflammatory cellular infiltrate with formation of hair follicles and sweat glands in comparison with a healthy skin. According to the chemical analyses of *Chlorella vulgaris*, the antimicrobial material was defined as a phenolic compound having the following formula C$_{14}$H$_{17}$NO$_4$ and the suggested structure could be 2-(1-hydroxy-2-(4-hydroxyphenyl)-2-methoxyethyl-4-oxopentane nitrile.

**Keywords:** Chlorella vulgaris, Antifungal, Antibacterial, Impetigo, Dermatophytes

Infectious diseases are caused by microbes including bacteria, fungi, protozoa, and viruses (WHO, 2010). Synthetic drugs are not only expensive and inadequate for the treatment of diseases but are also often with adulterations and side effects. Therefore, there is a need to search for new infection-combating strategies to control microbial infections (Sieradzki et al., 1999). Pharmaceutical industries are increasingly recognizing the importance of compounds derived from soil plants and other sources such as marine organisms (McGee, 2006).

Algae are a source of amino acids, terpenoids, phlorotannins, steroids, phenolic compounds, halogenated ketones, alkenes and cyclic polysulphides (Taskin et al., 2007). A large number of algal extract products have been found to have antimicrobial activity, many of the structures were identified as fatty acids and hydroxyl unsaturated fatty acids, glycolipid, steroid, phenolics and terpenoids.

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Lauric acid, palmitic acid, linolenic acid, oleic acid, stearic acids are known to be potential antibiotic or antifungal agents (Tan, 2007).

All of humans are colonized by more bacterial cells than human cells they have in their bodies. Generally, this is a peaceful and even productive (symbiotic) relationship, but occasionally even these well-tolerated residents of the human biosphere cause disease (Relman, 2002). Raman et al. (2005) tested chloroform-methanol mixture, n-butanol and diethyl ether extracts of two marine brown algae (Sargassum vulgare and Padina tetrasormatica) from Bay of Bengal near Visakhapatnam sea coast and evaluated for antibacterial activity by agar-well diffusion method. The diethyl ether and chloroform-methanol extracts of S. vulgare and diethyl ether, chloroform-methanol and n-butanol extracts of P. tetrasromatica exhibited significant antibacterial activity against the 4 tested bacteria namely Escherichia coli, Klebsiella pneumoniae, Bacillus subtilis and Staphylococcus aureus. Moreover, antibacterial activity of methanolic extracts from 20 species of macroalgae (9 Chlorophyta, 3 Phaeophyta and 8 Rhodophyta) collected from Moroccan Mediterranean coasts was evaluated against E. coli, S. aureus and Enterococcus faecalis (Zbakh et al., 2012). The extracts of the studied Rhodophyceae inhibited considerably the growth of the three tested bacterial strains. The results indicated that these species of seaweed have a significant capacity of antibacterial activities, which makes them interesting for screening for natural products (Zbakh et al., 2012).

Dermatophytes are a highly specialized group of fungi that, through long evolutionary processes, became adapted to invade, colonize and nourish themselves on the keratinized tissues of animals. Many investigators have shown that these diseases producing moulds possess enzymatic systems that enable them to digest keratins (Weary and Canby, 1969). They affect 20% to 25% of the world’s population, and the incidence continues to increase (Havlickova et al. 2008). Most cases of tinea unguium, tinea cruris, tinea corporis, and tinea pedis are caused by Tinea rubrum, which is the most common dermatophyte in developed countries (Borman et al., 2007). The phenolic compounds released from dried crudes of seaweed extracts may be answerable for their antimicrobial properties. This was confirmed earlier by Cox et al. (2010) who found that phenolic compounds are responsible for the antifungal activities of seaweeds. This may be due to the impact of these antifungal compounds on spore germination (El-Mehalawy, 2003).

This work aims to study the in vivo efficiency of Chlorella vulgaris methanolic extract to be used in the treatment of impetigo, tinea corporis and cutaneous candidiasis infections.

Materials and Methods

Alga and Growth conditions

Chlorella vulgaris (Chlorophyta) was obtained from the culture collection of Botany Department, Faculty of Science, Mansoura University, Egypt. Kuhl’s medium (Kuhl, 1962) was used for cultivation of C. vulgaris. The culture was aerated with air mixed with 3% CO₂ and incubated at 28 °C under continuous
illumination provided from day light fluorescent tubes giving light intensity of 80 \( \mu \text{E m}^{-2} \text{s}^{-1} \). *Chlorella vulgaris* was grown and harvested on 12\(^{th}\) day. The collected biomass was dried in an oven at 40-60°C and then powdered by a mortar.

**Algal extracts**

The algal powder of *C. vulgaris* was soaked in 70% methanol for 5h at room temperature and sonicated for 15 min. algal extract was filtered through Whattman filter paper No.1. The obtained filtrate was freed from solvent by evaporation under reduced pressure. The obtained residues (crude extracts) were suspended in methanol to a final concentration of 50 mg/ml. The extract was stored in the refrigerator in airtight glass bottle for further experiments.

**Isolation and culture techniques of pathogenic microorganisms**

Samples were collected from patients clinically diagnosed to have impetigo, tinea corporis and cutaneous candidiasis under aseptic conditions. Prior to sampling, the affected area surface was cleaned with cotton swab moistened with 70% ethyl alcohol. Different parts on mouse, near nose, arms and legs (for impetigo samples), legs and arms (for tinea corporis and cutaneous candidiasis samples), cotton swabs, tubes of phosphate buffered saline, glass slides and Petri dishes were previously sterilized to be used in sampling. Sterile scalpel was used to collect skin scrapings on sterile glass slide (El-Shanawany, 1993). The collected samples were isolated in a closed controlled room in hospital laboratory.

**For the impetigo samples**

Samples were forwarded to the Bacteriology Laboratory in 2 ml phosphate-buffered saline (PBS) composed of NaCl, 8 g/L; KCl, 0.2 g/L; Na\(_2\)HPO\(_4\), 1.15 g/L; and KH\(_2\)PO\(_4\), 0.2 g/L. All culture swabs were processed in the same day that they were collected. Each specimen was plated to mannitol salt agar medium. Culture plates were incubated up to 24 hr at 37°C (Gheda *et al*., 2012).

**For the tinea corporis and cutaneous candidiasis samples**

Each sample was cultured in three sterilized Petri dishes containing sterile Sabouraud's dextrose agar (SDA) medium; each 1 liter of distilled water contained 30 g dextrose, 10 g peptone, 20 g agar, and 0.5 g from each chloramphenicol and cycloheximide were added to avoid bacterial contamination (Al Doory, 1980). Plates were incubated at 37°C for 7-20 days for dermatophytes and for 2-3 days for *Candida albicans*.

**Treatment of impetigo and dermatophytes in experimental animal**

Healthy albino rats of average weight of 120 g were processed and placed in animals' house, Zoology department, Faculty of Science, Tanta University. Fresh isolates of *Staphylococcus aureus, Microsporum canis* and *Candida albicans* were utilized to perform artificial infections with the pathogens. The rats were infected with the different pathogens after producing the thermal lesion on its back as follows: after anaesthetization of the animal, a pre-heated brass blocks (92–95 °C) to the backs of the shaved rate for 5 sec. After the infliction of burns, the eschars were immediately injected intradermally with \(10^6\) CFU of all pathogens (Stevens *et al*., 1994). The infection repeated daily and the rats were
left a time to allow the infection to form a definite lesion which appeared after 5 days for bacteria and after two weeks for fungi.

*Chlorella vulgaris* ointment was prepared from its extract and the topical cream was carried out according to Purushothamanrao *et al.* (2010). The prepared cream was applied on the lesion separately twice a day (5 rats for cream treatment) for a total period of one week for the bacterial infection and two weeks for the fungal infection. Positive control of ciclopirox and nystatin creams were applied for 5 infected rats as antifungal agent for 5 rats, while mupirocin was applied as antibacterial for another 5 rats. Also, negative control was conducted by leaving the infection lesion without treating.

**Histopathological examinations of rat’s skin tissues**

Histopathological examination was carried out for 4 groups of rats. The first one was for healthy rat’s skin tissue as a positive control. The second was for infected rat’s skin tissues with impetigo and dermatophytes (untreated) as a negative control. The third was for infected rat’s skin tissues treated with algal extract ointment. The last group was for infected rat’s skin tissues treated with mupirocin, ciclopirox and nystatin. Skins were cut using a sterile surgical blade and immediately soaked in 10% formaldehyde fixative solution, and left for 24 hr. Skin samples were soaked in 95% ethanol with traces of eosin dye to be distinguished, then clarified by soaking in xylene (miscible with paraffin) to avoid turbidity of ethanol. Skin samples were transferred into molten soft paraffin bath. Sections of 5µm thickness were made from paraffin blocks containing skin samples by rotary microtome with clean sharp heated biconcave knife (Alcon-Couvreur, Belgium), and then the skin sections were stained. The skin sections were examined using light microscope, sharp images were photographed with magnification power of (200x and/or 400x).

**Chemical analysis of *C. vulgaris* extract**

**Determination of the chemical structure of antimicrobial material**

A.  Mass spectra (MS) of the antimicrobial material

A mass spectrophotometer (Shimadzu Qu-2010 Plus) was used for subjection of the compound to a stream of high energy of electrons at elevated temperature up to 100 °C. The produced fragments were yielded which can be characterized by mass/charge from spectra data.

B.  Proton nuclear magnetic resonance (1H NMR) spectra

The sample was dissolved in dimethyl sulfoxide (DMSO). The different protons of functional groups were identified using NMR (Varian Gemini 200 MHZ).

**Results**

Albino rats were infected with fresh isolates of *Staphylococcus aureus*, *Microsporum canis* and *Candida albicans*. The inoculation area was shaved as in photos (Photo 1.a) and burned as shown in photo (Photo 1.b). The infection began on the stratum corneum by allowing contact of pathogens cells of *S. Egypt. J. Bot.*, 56, No. 2 (2016)
inoculation of impetigo, tinea corporis and candidiasis appeared. The symptoms of scaly lesions were observed with inflammation appeared at the infected area as in (Photos 1.c1, 2, 3). After complete establishment of the lesions, the following treatments were applied; first, ointment preparation of C. vulgaris extract that possesses a high inhibitory effect on S. aureus, M. canis and C. albicans was applied separately on the infected lesions of rats skin. Second, mupirocin, ciclopirox and nystatin creams were applied separately on the infected lesions of other infected rats. Sections from the rats under treatment showed the healing process and examining the changes in the layers of skin.

Photo 1. In vivo healing of Staphylococcus aureus, Microsporum canis and Candida albicans growth in rat’s impetigo, tinea corporis and candidiasis lesions at different stages: Control: a: Non-infected, non-treated rat skin, b: Non-infected, non-treated, burned rat skin, c: Infected, non-treated rat skin with high occurrence of lesions 1- S. aureus 2- M. canis 3- C. albicans d: Infected rat skin after 4 days of treatment, 2- S. aureus 2- M. canis 3- C. albicans

The treatment was very effective after one week of treatment of impetigo as shown in Photo 1.e1 and partial healing of skin infected by fungi appeared as shown in Photos 1.e2, 3, while after two weeks the hairs appeared again in the rats infected with bacteria as shown in Photo 1.f1 and the complete healing of the fungal infections appeared as shown in Photos 1.f 2, 3. Our study revealed that the treatment with mupirocin, ciclopirox and nystatin after one and two weeks, respectively, illustrates partially cure of lesions and inflammation.

**Photo 1 Cont . e: Infected rat skin after 1 week of treatment,**
1- S. aureus 2- M. canis 3- C. albicans

**f: Infected rat skin after 2 weeks of treatment,**
2- S. aureus 2- M. canis 3- C. albicans

Improvement tests for C. vulgaris extract effectiveness and safety to be used in topical treatment for impetigo and fungal infections were achieved by histopathological studies (Photo 2). Skin sections for healthy rat skin tissues showed normal histological pattern; normal epidermis (Ep) as the keratinized fibres of stratum corneum (→) were regularly arranged, appeared condensed without any disruption and the dermis (D) appeared normal which formed of two layers; papillary and reticular layer with normal fibroplasts, hair follicles, sebaceous glands in papillary layer and sweat glands. The reticulate layer appeared with many fat cells as shown in (Photo 2.a). Skin tissues of the infected rats (untreated) showed; in case of the bacterial infection, partial losing of the epidermis layer with lost, disrupted stratum corneum layer (→) and the dermis showed inflammatory cellular infiltrates (*) mainly formed of lymphocytes and plasma cells as shown in (Photo 2.b.1). However, in case of the fungal infections,
the infection caused by *Microsporum canis* showed scap (S) covering the upper part of lesion, and the thickness of the underling epithelium is apparently smaller (→) than the epithelial layer of the normal part in the same section as shown in (Photo 2.b.2). The infection caused by *Candida albicans* showed abnormal histological pattern of skin; as the dermis showed infiltration (*) and loss of the hair follicles (?) as shown in (Photo 2.b.3).

Photo 2. Histopathological effects of Chlorella vulgaris extract treatment for impetigo, tineacorpsis and candidiasis infections on rat’s skin lesions against mupirocin, ciclopirox and nystatin;  

- a- Negative control: healthy skin,  
- b- Positive control: infected non-treated skin,  

1-S. aureus 2- M. canis 3- C. albicans

During the period of treatment, the healing of skin appeared gradually and the histological effects of *C. vulgaris* extract treatment were followed up periodically. The skin returned to its normal pattern in a short time and the skin

tissues appeared with normal epidermis as the keratinized fibres of stratum corneum were regularly arranged (→) appeared condensed without any disruption and the dermis appeared normal with minimal inflammatory cellular infiltrate (*) as shown in (Photo 2.c). On the other hand, mupirocin-treated skin sections showed normal epidermis with apparently normal thickness, the thickness of stratum corneum layer appeared normal in a part of section but appeared thin (→) in another part in the same section. Regarding to the dermis; it still loses the hair follicles (?) as shown in (Photo 2.d). Ciclopirox- and nystatin-treated skin section possessed stratum corneum was still lost (→), separation of epidermis and the dermis (►) showed exudates with some inflammatory cellular infiltrate of lymphocytes, vacuolation of keratinocyte or epidermal cells (*) as shown in (Photo 2.e).

Photo 2. (Cont.) Histopathological effects of *Chlorella vulgaris* extract treatment for impetigo, tinea corporis and candidiasis infections on rat's skin lesions against mupirocin, ciclopirox and nystatin;

a. *C. vulgaris*-treated, healed skin,
b. Mupirocin-treated, partially healed skin,
c. Ciclopirox- and nystatin -treated, partially healed skin.

In the present study, NMR and Mass spectra data was used to elucidate the chemical structure of the antimicrobial material extracted from *C. vulgaris*. The characteristic signals in the NMR spectrum were represented graphically in Fig. 1. The NMR spectrum showed two signals at δ 8.0 - 8.4 ppm which are characteristic of the 4 H aromatic protons. The two signals within δ 4.5 - 4.7 ppm range corresponding to 2 OH groups. The signals within δ 2.5 - 2.7 ppm range are characteristic of the protons of the CH group. The signals at δ 1.9 - 2 ppm region are characteristic of the protons of the OCH₃ group. The CH₃ group leads to a broad signal at δ 1.7 ppm with integration equivalent to one proton.

![Proton magnetic resonance of the antimicrobial material isolated from *Chlorella vulgaris*.](image)

The mass spectrum (MS) fragmentation pattern of the compound under investigation was shown in Fig. 2. It revealed the presence of peak at m/z 263 of relative abundance characteristic of the parent compound.

According to the data obtained, the antimicrobial material is a phenolic compound having the following formula C₁₄H₁₇NO₄ and the suggested structure should be 2-(1-hydroxy-2-(4-hydroxyphenyl)-2-methoxyethy 1-4-oxopentanenitrile. The mass spectroscopy of our antifungal material indicated that the molecular weight is 263 (Fig. 3).

Discussion

This study confirmed the high activity of *C. vulgaris* methanolic extract at low concentration. Results confirmed that it would treat the impetigo and skin fungal infections as Candidiasis and tinea corporis. Most researchers have attributed the degradation of keratinic substrates to the production of specific and mostly extracellular proteolytic enzymes called keratinases, whose secretion appears to be induced by the presence of keratin in the substrate (Apodaca and McKerrow, 1989; Siesenop and Böhm, 1995). However, those secreted by *T. rubrum* appear to be responsible for keratinolysis (Kwon-Chung and Bennet, *Egypt. J. Bot.*, 56, No. 2 (2016))
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Extracellular keratinases are produced by bacteria (Böckle et al., 1995; Lin et al., 1995) and fungi (Siesonop and Böhm, 1995; Lambkin et al., 1996). Roberts (1996) reported that the production of enzymes such as phospholipase, lipase and keratinase by the invading fungus could lead to damage of keratinocytes and affect melanocytes causing changes in skin structure. So, these enzymes may help in the passage of fungi to stratum corneum of skin by physical barriers and other layers of skin and this could be a reason of tissue damage (Howard, 1983).

Virulence of pathogenic fungi depending on keratinolytic activity was confirmed by analysis of virulence factors (Viani et al., 2001) as fungal pathogenesis was proved by keratinase production, which was characterized and purified from Microsporum canis. Many filamentous fungi can synthesize a diverse range of hydrolytic enzymes such as proteases and lipases (Weitzman and Summerbell, 1995). However, keratinases are the key enzymes in fungal invasion of skin and have been mostly studied in dermatophytes, including Trichophyton sp., Microsporum sp., some pathogenic yeasts such as C. albicans and some other fungi and bacteria (Okafur and Ada, 2000; Muhsin and Hadi, 2002). Gheda et al. (2012) tested the antibacterial activity of Spirulina platensis for the treatment of patients infected by impetigo, and they reported that in vivo application of both active ingredients and crude extracts of Spirulina platensis showed promising response rates, and no side effects appeared during the follow-up period.

The chemical analyses of algal extract by methanol showed the presence of active phenolic compounds. Vepritski et al. (1991) also reported that cyanobacterin LU-2 produced by Nostoc sp. is a phenolic derivative containing amino-sugar. The produced antimicrobial material is effective against fungi, which was agreed with that reported by De cano et al. (1990) who observed inhibition of C. albicans by phenolic compounds from N. muscorum and Chlorella sp. The mass spectroscopy of our antimicrobial material indicated that the molecular weight is 263. Glombitza and Damhues (1985) studied the chemical structure of 17 oligomeric chloro-tannins isolated from the brown alga Himanthalia elongate using 1H NMR and mass spectra. Sastry and Rao (1994) identified the antimicrobial compound isolated from S. wightii as dioctyl phthalate. The structure was confirmed from spectroscopic data ($^1$H and $^{13}$C NMR), which were compared with data obtained from authentic samples. Jaki et al. (2000) determined the structure of five novel extracellular di-terpenoids with biological activity from the cyanobacterium N. commune by spectroscopic methods, mainly NMR and MR. Furthermore, Jaki et al. (2001) determined the structure of two novel cyclic peptides with antifungal activity from the cyanobacterium Tolypothrix by 1D and 2D NMR experiments and tandem mass spectrometry.

Conclusion

The results of this investigation showed that the methanol extract of C. vulgaris used as natural ointment was effective against bacterial and fungal skin infections. The histological examinations of rat's skin layers revealed the fast healing of the skin layers and showed normal histological patterns of rat skin, while the synthetic ointments showed partial healing comparing with the natural

product from *C. vulgaris* extract ointment. The chemical analysis suggested the chemical formula of the antifungal substance as $\text{C}_{14}\text{H}_{17}\text{NO}_4$ and the suggested structure should be 2-(1-hydroxy-2-(4-hydroxyphenyl)-2-methoxyethyl-4-oxopentanenitrile.

References


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IN VIVO EVALUATION OF ANTIMICROBIAL EFFECT FROM THE MEAN CRYSTAL METHANOLIC EXTRACT OF SOME PLANTS

Some plants, such as Spirulina platensis, have been used to treat some diseases, such as mycosis, which is a condition caused by fungi. The mucor fungi are a group of fungi that cause diseases such as mycosis. The study aims to evaluate the antimicrobial activity of the mean crystal methanolic extract of some plants against a number of pathogenic fungi, including Mucor. The results showed that the extract had a significant antimicrobial effect against the tested fungi. The study also aimed to evaluate the effects of adding some amino acids to the extract to improve its effectiveness. The results showed that adding some amino acids to the extract improved its antimicrobial activity. 