SPONGE-ASSOCIATED FUNGI ISOLATES FROM ANCORINA SP. SHOWED ANTI-CANCER ACTIVITY AGAINST HELa CELL LINES

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ABSTRACT

Nowadays, natural products from marine life, especially sponge-associated fungi (SAF), have been getting extensive attention as candidates for anti-cancer agents. Although chemotherapy has been widely used as one of the cancer treatments, recently, multidrug resistance as one of its drawbacks has appeared. Our preliminary research successfully isolated 16 fungi isolates from Ancorina sp. inhabiting Kukup (KU) Beach, Yogyakarta, Indonesia. The ethyl acetate extracts of SAF KU4 fungi grown in enrichment broth media in seawater base either as media-free mycelium or whole medium were analyzed for the cytotoxicity level and the apoptotic effect on the apoptotic effect cervical cancer cell line (HeLa). SAF KU4 was cultured in Wickerham sea-based media and then extracted using ethyl acetate. The IC₅₀ values were measured by MTT assay. IC₅₀ of the media-free extract and whole extract were 158.13 and 283.95 µg/mL, respectively. Ethyl acetate extracts of media-free mycelium and whole extract were moderately cytotoxic but could induce apoptosis in HeLa cell lines.

Keywords: Ancorina sp., Apoptosis, Cytotoxicity, HeLa cell, SAF KU4

INTRODUCTION

Cancer is still one of the leading health problems and is getting more attention in biomedical research, especially in obtaining new compounds with anti-cancer activity. Furthermore, the cause of cancer and its manifestation are still unclear and need to be elucidated (Beesoo et al., 2014; Dey et al., 2015). Chemotherapy is an optional treatment for cancer patients since it is well-documented that chemotherapeutic drugs induce multiple-drug resistance in cancer cells. Chemotherapy drugs produced by synthetic chemistry have become a concern since these drugs are lethal to both cancer and healthy cells; thus, interest in natural products as anticancer has increased recently (Haufler, 2003; Aung et al., 2017). Natural products isolated from fungi are promising since the first isolation of cyclosporin A from Tolypocladium inflatum was approved for clinical treatments as an immunosuppressant in 1983 (Bugni & Ireland, 2004). Meanwhile, the high rediscovery of isolated compounds has interested researchers in exploring unique habitats such as the marine environment. Microbes, i.e., fungi, alive in this environment have constantly been exposed to extreme conditions, like temperature, salinity, pressure, etc., that can produce various compounds with unique structure and biological activity (Bugni & Ireland, 2004; de Carvalho and Fernandes, 2010; Deshmukh et al., 2018). Sponge-associated fungi produce natural products with active substances showing antibiotic, anti-helminthic, anti-neoplastic, and anti-cancer activities. For instance, Neochelinula A and Physicin, two active compounds isolated from Microsporum sp., and algae-associated fungi, Lomentaria catenata, are known to have anti-cancer activity and are being developed as new potential candidates for chemotherapy (Wijesekara et al., 2013, 2014).

Up to date, the exploration of novel compounds from sponge-associated fungi is fascinating and has the potency to identify new drugs for cancer. This study isolated 16 sponge-associated fungi from Ancorina sp. inhabiting Kukup Beach, Yogyakarta, Indonesia, and screened them for anticancer properties. Here we report the anti-cancer activity of an Aspergillus named SAF KU4, one of the 16 isolates that showed anti-cancer activity with the lowest IC₅₀ tested against HeLa cells. Further, we also standardized the conditions that enhance the production of these anti-cancer active compounds from SAF KU4.

MATERIAL AND METHODS

Materials

HeLa cell lines were from LPPT (Laboratorium Penelitian dan Pengujian Terpadu/The Integrated Research and Testing Laboratory, Universitas Gadjah Mada. SAF KU4 stock isolate was stored at 4°C in the slant of malt extract agar. Dimethylsulfoxide (DMSO) (NacalaTeseq), 3-(±)-dimethyl-2-yl)-2,5-diphenyloxazole (MTT) (Sigma-Aldrich), RPMI 1640 (Sigma-Aldrich), Fetal Bovine Serum (Sigma-Aldrich), Penicillin-Streptomycin (Sigma-Aldrich), Fungizone (Gibco), Phosphate Buffer Saline, Stop solution (10% SDS in 0.01N HCl), Acridine Orange (Sigma-Aldrich)/Ethidium Bromide (Sigma-Aldrich), ethyl acetate (EtOAc), malt extract agar (Sigma-Aldrich), monohydrate glucose (Sigma-Aldrich), peptone (Oxoid), and yeast extract (Oxoid).

Subculture and identification of SAF KU4

SAF KU4 was isolated from the sponge, Ancorina sp., inhabiting sea rocks at Kukup Beach, Yogyakarta, Indonesia in 2014. For preculture, SAF KU4 was grown in Malt Extract Agar (15 g of agar, 15 g of malt extract, 1000 mL of seawater). Identification was proceeded by the lactophenol staining. A 10-day old culture of SAF KU4 fungi with hyphae was fixed to object glass and dropped by lactophenol (Sigma-Aldrich). Furthermore, the structure hyphae, spore shape, mycelia turbidity, and mycelial branching were observed under the light microscope (Olympus) with a maximal magnification of 1,000x. The pictures were taken by a camera (Sony Camera).

Cultivation of SAF KU4 and secondary metabolite extraction

Two flasks, each containing 300 mL Wickerham Medium (5 g of yeast extract, 3 g of malt extract, 5 g of peptone, 20 g of monohydrate glucose, 1000 mL of seawater) were used to inoculate 3% (v/v) of SAF KU4 fungus and cultivated for 21 days at room temperature with agitation at 200 rpm. After 21 days, one flask was extracted after removing mycelia (media-free mycelia) with ethyl acetate (EtOAc) (1:1 v/v), and these were macerated for seven days (Noimart et al., 2017). The ethyl acetate phase was separated from broth and evaporated until obtaining the media extract. Another flask containing the mycelia in the medium was disrupted by ultrasonication, and the lysate was filtered by Whatman Paper No. 1 in Vacuum Buchner funnel. The filtrate devoid of cell debris was extracted separately with
Repeat the process of extending the aseptate conidiophore and modified into C-ration, cells were treated with vesicles in Figures 2(A) and 2(B). Using the probit-50 method, the cell was enriched Wickerham media with prolonged incubation until 21 days. We then obtained the cytotoxicity curve of both extracts using extract concentrations for analysis from 600-1.7 µg/mL. The whole extract showed about 61% of HeLa cell line was apoptotic at 200 µg/mL, and it showed a decrease in cell viability by about 51% at IC-50. Probit analysis was conducted to determine the IC-50 of both two extracts. The number of apoptotic and healthy cells in the different extracts were compared to two times concentration higher from the IC-50 concentration values. The effect of two extract treatments in HeLa cell lines is shown in Figure 3. Several variations in apoptotic and healthy cells were observed between media and combination extract treatments. In addition, lower concentration from the IC-50 value showed a high number of healthy cells compared to two times concentration higher from the IC-50 value.

RESULTS AND DISCUSSION

Morphology of SAF KU4
Fungi can be determined by observing the hypha and conidiophore structures under the microscope. We assumed SAF KU4 was classified into the Ascomycota class in the genus Aspergillus based on the morphological observation. This genus has a foot cell produced by an asci and conidiophore and modified into vesicles (Bennet, 2010; Samson, 1994). SAF KU4 was first isolated from the sponge Ancorina sp. (Figure 1A) inhabiting Kukup beach, Yogyakarta, Indonesia. The sponge is found at the intertidal zone attaching to the shoreline rocks. The microscopic structure of the fungi under the microscope showed a fungal structure from the Aspergillus genus (Figures 1B & 1C). SAF KU4 has modified hyphae as a vesicle and aseptate conidiophore with transparent unbranched hyphae. Under the microscopic structure analysis, SAF KU4 has an aspergillom-like spore-bearing structure which is a key character to identifying Aspergillus genus accurately.

Figure 1 Morphology of sponge Ancorina sp. from Kukup beach, Yogyakarta, Indonesia (A); Microscopic morphology of SAF KU4 (B and C) isolated from sponge Ancorina sp. vesicles (black rectangle, 400x) and aseptate conidiophores (red rectangle, 1000x).

Extraction and Cytotoxicity Assay of SAF KU4 Secondary Metabolites
We successfully obtained 0.18 g extract of both whole and media extracts, which showed an oily appearance with a brownish color. Then, assays of two extracts were continued against HeLa cancer cell lines. The cytotoxicity assay was determined by MTT method. Our preliminary research revealed that SAF KU4 extract, cultured in malt extract with 14 days incubation, showed IC-50 around 300 µg/mL. This value was then referred to assay SAF KU4 extract cultured in enrichment Wickerham media with prolonged incubation until 21 days. We then used extract concentrations for analysis from 600-1.7 µg/mL. Based on the assays, we obtained the cytotoxicity curves in Figures 2(A) and 2(B). Using the probit analysis, MTT assay on HeLa cell lines showed the IC-50 of media extract and whole extract were 158.13 and 283.95 µg/mL, respectively.

Figure 2 Effect of media extract (A) and combination extract (B) on HeLa cancer cell lines.

Apoptosis Staining in HeLa cell lines
The Acridine Orange/Ethidium Bromide method was used to distinguish between healthy cells and apoptotic cells. Here, HeLa cell lines were treated using two times lower and higher IC-50 concentration values. The effect of two extract treatments in HeLa cell lines is shown in Figure 3. Several variations in apoptotic and healthy cells were observed between media and combination extract treatments. In addition, lower concentration from the IC-50 value showed a high number of healthy cells compared to two times concentration higher from the IC-50 value.

Figure 3 HeLa cell lines condition in acridine orange/ethidium bromide staining after treatments. A) control; B) media extract at concentration 85 µg/mL; C) media extract at concentration 170 µg/mL; D) whole extract at concentration 100 µg/mL; E) whole extract at concentration 200 µg/mL. Observation took under a fluorescent microscope (40x). The white arrow shows the healthy cell, and the yellow one is the apoptotic cell.

In Figure 4, the number of apoptotic and healthy cells in the different extracts were shown. Media extract with a concentration 170 µg/mL showed about 61% HeLa cell line in apoptotic condition, and then, it was decreased to 56% at 85 µg/mL. The whole extract showed about 61% of HeLa cell line was apoptotic at 200 µg/mL, and it showed a decrease in the number of apoptotic cells by about 51% at 100 µg/mL.
**Figure 4** The percentage of HeLa cells in the healthy condition (black) and apoptotic condition (grey) after treatment of media extract (A) and whole extract (B). It could be observed that apoptotic cells increased in a high concentration of extracts.

Sponge extracts of Ancorina sp. were previously reported to have anti-cancer activity (Nuriliani et al., 2013; Tunjung & Sayekti, 2019), but the activity of sponge-associated fungi extract is not reported well. Here, we first reported isolated sponge-associated fungi from Ancorina sp. that showed anti-cancer activity (Ramadhani et al., 2017). Previously, we successfully isolated 16 isolates of fungi from the sponge living at Kukup beach. Antibacterial activity against Staphylococcus aureus and Salmonella typhi was used as preliminary method to observe the antibiotic activity. Two candidate isolates, SAF KU3A and SAF KU4, showed the highest antibiotic activity in bacterial growth inhibition from sixteen isolates. We predicted that we would have promising activity as the anti-cancer agent. Further analysis, by using cervical cancer cell line HeLa, ethyl acetate extract from SAF KU4 showed the highest cytotoxicity value compared to SAF KU3A (Ramadhani et al., 2017), which then, was chosen as the candidate to investigate the activity against HeLa cancer cell lines with modification in cultivation medium, prolonged the cultivation time, and added agitation process. Filamentous fungi are known as the source of bioactive compounds (Chavez et al., 2015). As the antibiotic story, Penicillin, the first well-known antibiotic from a microorganism, is isolated from fungi known as Penicillium notatum. Meanwhile, recently the number of resistance problems, even for the bacteria (antibiotic) and cancer cell (anti-cancer), has increased periodically, demanding new drugs needed to be approved by the authorities. The attempt to harness fungal secondary metabolites for drug discovery gives an approach to isolating new compounds from new habitats of isolation, such as fungal associated with a sponge. Sponge-associated fungi have been known to produce novel and unique secondary metabolites (Tian et al., 2018; Zhou et al., 2011). The interaction of fungi and sponge is mediated by a specific site (1→3)-β-D-glucan binding protein. This protein is in the sponge surface and makes a compatible site for fungi to attach to the sponge body and being associated (Suryanarayanan, 2012). This binding protein is also hypothetically synthesized and located on the surface of Ancorina sp., which makes a possible interaction with SAF KU4.

In the first preliminary research, we obtained the IC50 higher than 300 µg/mL against HeLa cell lines (Ramadhani et al., 2017). It was known that modifying and adding substances in growth media would affect active metabolite production (Miao et al., 2006). Increasing activity and production of active compounds from a microorganism, especially from fungi, can be engineered by modification of incubation time (Frisvad et al., 2007), stress given during culture, i.e., salt concentration (Wang et al., 2011), or modification on a culture medium (Marmann et al., 2014). In our studies, malt extract was used for cultivation media at the preliminary stage. It was also reported that enrichment of carbon, hydrogen, oxygen and nitrogen sources in cultivation medium raised the secondary metabolite production and mass of fungi (Yang et al., 2007). Carbon, hydrogen, and oxygen are the primary backbone of secondary metabolite scaffold, and nitrogen can be found occasionally on the compound and makes unique structures to be produced (Nursid et al., 2010). Further, we tried to modify the medium for the culture by using Wickerham media as the enrichment media, and we added agitation. Aeration on liquid culture is proven to actively increase the fungi biomass and increase the production of secondary metabolites. Aspergillus, the reported secondary metabolite produced by Aspergillus glaucus HB 1-19, showed an increase in production after being given the agitation at 90 rpm (Cai et al., 2012). Therefore, maintaining SAF KU4 with agitation can further increase the production of active secondary metabolites.

Production of secondary metabolites mostly occurs at the stationary phase due to a lack of nutrients; thus, the microorganisms compete to get any available remaining nutrients from the environment (Calvo et al., 2002). Those secondary metabolites can either be accumulated inside or excreted outside of the cells, which depends on the function of the cell. If toxic compounds are produced inside the cell, they will be passed outside the media and protect the cell from death, making cells alive and getting nutrients. However, the favorable compound was still maintained inside the cell and used as a growth substance (Yining, 1990). The presence of active compounds from SAF KU4 was investigated by comparing the extract of media and whole extract by assaying the bioactivity of the secondary metabolites from extracellular and the combination of extracellular and intracellular compounds. The results showed that ethyl acetate extract from media gave the lowest IC50 compared to the whole extract, and it still had a potency as the anti-cancer agent. Meanwhile, the IC50 from mycelium extract was reported previously at 164 µg/mL (Ramadhani et al., 2017). This IC50 showed comparable cytotoxicity compared to media and the highest cytotoxicity compared to the whole extract. Here, we suspect that the active compounds from SAF KU4 were still located outside of the cells. However, we also predict some active compounds are also still located inside the cells, proven by the comparable IC50 between media and mycelium ethyl acetate extracts. The whole extract showed the highest IC50 compared to media and mycelia extracts. Yet, it still showed the lowest IC50 value compared to the results from the preliminary research with an IC50 value of 383.88 µg/mL. We assume that compound-compound interaction may be the main reason the IC50 value from the whole extract is highest compared to medium and mycelium extracts. One study identified this mechanism and reported that compounds could interact with each other to make antagonistic effects on cells that will change the ability of active compounds to inhibit the growth of the targeted cells (Yin et al., 2014). Meanwhile, in our research, we found that modification on culture medium, prolonged incubation time, and agitation process during the culture process changed the IC50 value of the SAF KU4 extract.

Apoptotic induction staining confirmed the presence of active compounds obtained by both extracts to induce the apoptotic body (pointed out by red/orange color, which the healthy cells are stained green color) (Figure 2). We found that both media and whole extracts showed increasing apoptotic cell body dose-dependence, suggesting their potency as an anti-cancer agent. It was known that ethyl acetate, a semi-polar solvent, could extract the active compounds, such as alkaloids, flavonoids, terpenoids, peptides, and other substances (Mandal et al., 2015). We suggest alkaloids, flavonoids, terpenoids, and peptides are present in both extracts and may induce the apoptotic in the HeLa cell line. However, further investigations are still needed to analyze the chemical compounds in both extracts. In addition, the pathway mechanism of apoptosis induction in the HeLa cell line by SAF KU4 extract also needs to be further investigated.

**CONCLUSIONS**

The modifications on media by enriching the nutrients, giving a prolonged day incubation, and adding agitation increased secondary metabolites production by SAF KU4 fungi isolated from the marine sponge. The ethyl acetate extracts of SAF KU4 induce apoptosis in HeLa cancer cell lines. The IC50 of media and whole extracts were 158.13 and 283.95 µg/mL, respectively.

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**REFERENCES**

Aung, T.N., Qu, Z., Kortschak, R.D., and Adelson, D.L. (2017). Understanding the effectiveness of natural compound mixtures in cancer through their molecular mode of action. Int. J. Mol. Sci., 18(3): 656. https://doi.org/10.3390/ijms18030656.

Beesoo, R., Neergheen-Bhujun, V., and Bhagooli, R. (2014). Apoptosis inducing lead compounds isolated from marine organisms of potential relevance in cancer treatment. Mutat. Res. Mol. Mech. Mutagen., 768: 84-97. http://dx.doi.org/10.1016/j.mrmmm.2014.03.005.

Bennet, J.W. (2010). An overview of the genus Aspergillus. In: Aspergillus Molecular Biology and Genomics, Machida, M. and K. Gomi, (Eds.). Caister, Academic Press, pp: 1-17.

Bugni, T.S., and Ireland, C.M. (2004). Marine-derived fungi: a chemically and biologically diverse group of microorganisms. Nat. Prod. Rev., 21(1): 143-163. http://dx.doi.org/10.1080/096708403901926h.

Cai, M.; Zhou, X-L.; Fan, W.; Zhou, J.; Niu, C.; Kang, L.; Sun, X.; and Zhang, Y. (2012). An integrated control strategy for the fermentation of the marine-derived fungus Aspergillus glaucus for the production of anti-cancer polyketide. Mar. Biotechnol., 14(6): 665-671. http://dx.doi.org/10.1007/s10292-012-9435-6.

Calvo, A.M., Wilson, R.A., Bok, J.W., and Keller, N.P. (2002). Relationship between secondary metabolism and fungal development. Microbiol. Mol. Biol. Rev., 66(3): 447-459. http://dx.doi.org/10.1128/MMBR.66.3.447-459.2002.
Chavez, R., Fierro,F., Garca-Rico,R.O., and Vaca,I. (2015). Filamentous fungi from extreme environments as a promising source of novel bioactive secondary metabolites. *Front. Microbiol.*, 6:1-7. http://dx.doi.org/10.3389/fmicb.2015.00903.

Deshmukh, S.K., Prakash, V., and Ranjan, N. (2015). Marine fungi: A source of potential anticancer compounds. *Front. Microbiol.*, 8:2536. http://dx.doi.org/10.3389/fmicb.2017.02536.

Dey, G., Bharti, R., Sen, R. and Mandal, M. (2015). Microbial amphiphiles: a class of promising new-generation anticancer agents. *Drug Discov. Today*, 20(1): 136-146. http://dx.doi.org/10.1016/j.drudis.2014.09.006.

Freitas, S., Lee, M., Silveira, M.A.L.N., and Wijayanti, N. (2013). Cytotoxic and apoptosis activity of sponge species of Astroporida order’s extract on HeLa cells (cervical cancer T47D cell line). http://dx.doi.org/10.3390/md15050139.

Frivsad, J.C., Larsen, T.O., de Vries, R., Meijer, M., Houbreken, J., Cabanes, F.J., Ehrlich, K., and Samson, R.A. (2007). Secondary metabolite profiling, growth profiles and other tools for species recognition and important Aspergillus mycotoxins. *Stud. Mycol.*, 59: 31-37. http://dx.doi.org/10.3114/SIM.2007.59.04.

Haefner, B. (2003). Drugs from the deep: marine natural products as drug candidates. *Drug Discov. Today*, 8(12): 536-544. http://dx.doi.org/10.1016/S1359-6446(03)00713-2.

Li, Y.-X., Himaya, S., and Kim, S.-K. (2013). Triterpenoids of marine origin as anticancer agents. *Molecules*, 18(7): 7886-7909. https://doi.org/10.3390/180500705.

Mandal, S.C., Mandal, V., and Das, A.K. (2015). Essentials of Botanical Extraction. Academic Press.

Marmann, A., Aly, A.H., Lin, W., Wang, B., and Proksch, P. (2014). Co-cultivation: a powerful emerging tool for enhancing the chemical diversity of microorganisms. *Mar. Drugs.*, 12(2): 1043-1065. http://dx.doi.org/10.3390/md12021043.

Miao, L., Kwong, T.F.N. and Qian, P.Y. (2006). Effect of culture conditions on mycelial growth, antibacterial activity, and metabolite profiles of the marine-derived fungus *Arthrinium* c.f. *saccariolica*. *Appl. Microbiol. Biotechnol.*, 72(5): 1063-1073. http://dx.doi.org/10.1007/s00253-006-0357-8.

Noinart, J., Buttcachon, S., Dethoup, T., Gales, L., Pereira, J.A., Urbatzka, R., Freitas, S., Lee, M., Silva, A.M.S., Pinto, M.M.M., Vasconcelos, V., and Kijjoa, A. (2017). A new ergosterol analog, a new bis-anthaquinone and anti-obesity activity of antimycotics from the sponge-associated fungus *Talaromyces stipitatus* KUFA 0207. *Mar. Drugs*, 15(15):139-151. https://doi.org/10.3390/md15050139.

Nurilani A., Ariyanto, I.A., Santi, M.R., Mahendra, A., Dewi, N.W.E.S., Huda, A.I.N., and Wijayanti, J., et al. (2013). Cytotoxic and apoptosis activity of sponge species *A. of Astroporida order’s extract on HeLa cells (cervical cancer cell line).* *Biota.* 18(1): 43-53. http://dx.doi.org/10.24002/biota.v18i1.263.

Nursid, M., Pratitis, A., and Chasanah, E. (2010). Kulitvisi Kapang MFW-01-08 Yang Disolasi Dari Ascidia Aplidium longithorax Dan Uji Aktivitas Sitotoksiknya Terhadap Sel Kanker Payudara T47D. *J. Pascapanen dan Bioteknol.* *Kelaut. dan Perikan.* 5(2):103-110.

Ramadhan, E., Priyambada, F., Pramana, A.A.C., Subchon, A.N., Pertivi, G.A., Setiawibawa, R.A.A., Putra, H.E., Amalia, N.R.A.S. and Wijayanti, N. (2017). Potential chemopreventive agent: study of apoptosis in the extracts of sponge-associated fungi from Yogyakarta against cervical cancer HeLa cell line. *In: International Conference on Applied Science and Health 2017: Improving Health and Well-Being for Better Society.* Mahidol University, 23 February 2017. ICASH., pp: 369-376.

Ribble, D., Goldstein, N.B., Norris, D.A., and Shellman, Y.G. (2005). A simple technique for quantifying apoptosis in 96-well plates. *BMC Biotechnol.* 5(1):1-12. http://dx.doi.org/10.1186/1472-6750-5-12.

Samson, R.A. (1994). Taxonomy: current concepts of Aspergillus systematics. *In: Aspergillus, J.E. Smith, (Eds.). Aspergillus.* Springer, pp: 1-22.

Suryanarayanan, T.S. (2012). The diversity and importance of fungi associated with marine sponges. *Bot. Mar.*, 55(6): 553-564. http://dx.doi.org/10.1515/bot-2011-0086.

Tian, Y.Q., Lin,S.T., Kumaravel,K., Zhou,H., Wang,S.Y. and Liu,Y.H. (2018). Polypeptide-derived metabolites from the sponge-derived fungus *Aspargillus* sp. F40. *Phytochem. Lett.*, 27: 74-77. http://dx.doi.org/10.1016/j.phytol.2018.06.009.

Tunjung, W.A.S, and Sayekti,P.R. (2019). Apoptosis induction on human breast cancer T47D cell line by extracts of *Ancorina* sp. from extreme environments as a promising source of novel bioactive secondary metabolites. *Front. Microbiol.*, 8:2536. http://dx.doi.org/10.3389/fmicb.2017.02536.

Vining, L.C. (1990). Functions of secondary metabolites. *Annu. Rev. Microbiol.* 44(1): 395-427. http://dx.doi.org/10.1146/annurev.mi.44.100190.002143.

Wang, Y., Lu, Z., Sun, K. and Zhu, W. (2011). Effects of high salt stress on secondary metabolite production in the marine-derived fungus *Spicaria elegans*. *Mar. Drugs*, 9(4): 535-542. http://dx.doi.org/10.3390/md9040535.

Yang, L.H., Miao, L., Lee, O.O., Li, X., Xiong, H., Pang, K.-L., Vrijmoed, L. and Qian, P.-Y. (2007). Effect of culture conditions on antifouling compound production of a sponge-associated fungus. *Appl. Microbiol. Biotechnol.*, 74(6): 1221-1231. http://dx.doi.org/10.1007/s00253-006-0780-0.

Yin, N., Ma, W., Pei, J., Ouay, Q., Tang, C. and Lai, L. (2014). Synergistic and antagonistic drug combinations depend on network topology. *PLoS ONE*, 9(4): 1-7. http://dx.doi.org/10.1371/journal.pone.0093960.

Zhou, Y., Mándi, A., Debbab, A., Wray, V., Schulz, B., Muller, W.E.G., Lin, W.-H., Proksch, P., Kurtan, T. and Aly, A.H. (2011). New ausstatlides from the sponge-associated fungus *Aspergillus* sp. *European J. Org. Chem.*, 6009-6019. http://dx.doi.org/10.1002/ejoc.201100670.