Conjugation of *Aspergillus flavipes* Taxol with Porphyrin Increases the Anticancer Activity of Taxol and Ameliorates Its Cytotoxic Effects

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Academic Editors: Bartosz Tylkowski, Anna Bajek, Krzysztof Roszkowski and Robert Musiol

Received: 24 November 2019; Accepted: 23 December 2019; Published: 9 January 2020

Abstract: Taxol is one of the potential anticancer drugs; however, the yield of Taxol and its cytotoxicity are common challenges. Thus, manipulating the Taxol biosynthetic pathway from endophytic fungi, in addition to chemical modification with biocompatible polymers, is the challenge. Four fungal isolates, namely, *Aspergillus flavipes*, *A. terreus*, *A. flavus*, and *A. parasiticus*, were selected from our previous study as potential Taxol producers, and their potency for Taxol production was evaluated in response to fluconazole and silver nitrate. A higher Taxol yield was reported in the cultures of *A. flavipes* (185 µg/L) and *A. terreus* (66 µg/L). With addition of fluconazole, the yield of Taxol was increased 1.8 and 1.2-fold for *A. flavipes* and *A. terreus*, respectively, confirming the inhibition of sterol biosynthesis and redirecting the geranyl phosphate pool to terpenoids synthesis. A significant inhibition of ergosterol biosynthesis by *A. flavipes* with addition of fluconazole was observed, correlating with the increase on Taxol yield. To increase the Taxol solubility and to reduce its cytotoxicity, Taxol was modified via chemical conjugation with porphyrin, and the degree of conjugation was checked from the Thin layer chromatography and UV spectral analysis. The antiproliferative activity of native and modified Taxol conjugates was evaluated; upon porphyrin conjugation, the activity of Taxol towards HepG2 was increased 1.5-fold, while its cytotoxicity to VERO cells was reduced 3-fold.

Keywords: Taxol; *Aspergillus flavipes*; fluconazole; porphyrin

1. Introduction

Taxol is a diterpenoid alkaloidal firstly isolated from *Taxus brevifolia* [1,2] as a strong antimitotic agent, causing cellular arrest at the G2/M phase of tumor cells. The anticancer activity of Taxol has been emphasized towards various tumor cell lines [3], including leukemia, breast, ovarian, and lung cancers [2,4], as well as against polycystic kidney diseases. Taxol production from different sources and their challenges have been addressed extensively on our studies [1,5,6]. Taxol is now being commercially produced by a semi-synthetic method, starting with 10-deacetylbaccatin III [7]...
extracted from the needles of *T. baccata* and *T. walichiana* growing in Europe and Asia. However, this method is challenged with lower yield, climatic dependence, and seasonal variations (reviewed by [5,8]). Chemical synthesis of Taxol has been emphasized [9,10]; however, the technical complexity and higher expenses are the main challenges stopping this technology from becoming an economically affordable approach [11,12].

Deciphering the potency of endophytic fungi for Taxol production opened a new avenue for commercial Taxol production due to their rapid growth, independence on climatic changes, and feasibility of genetic manipulation [5,13]. *Taxomyces andreanae* was the first recognized Taxol-producing endophyte of *Taxus* spp [14], after which an unlimited number of studies became motivated to isolate endophytic fungi with Taxol metabolic-producing potency [5–8]. Nevertheless, the implementation of these fungi for commercial production of Taxol has been confronted by the loss of Taxol productivity with subculturing and storage [1,6]. Manipulating the Taxol yield by specific inhibitors to modulate the metabolic yield by endophytic fungi was emphasized in our previous study [1]. Similarly, triggering the biosynthesis of secondary metabolites by plants in response to an external effector was also reported [15]. Additionally, using metabolic inhibitors to divert the metabolic flux toward the interested product has also been documented, especially the use of sterol biosynthesis inhibitors to divert the geranyl–geranyl pyrophosphate pool to Taxol biosynthesis, enhancing Taxol yields of up to 50-fold [16].

With the global commercial and clinical values of Taxol, the side effects and required high doses are still the major challenges to this drug, especially in developing countries. Conjugation with various biocompatible polymers is a reliable approach to increase the efficiency of this drug and to reduce its side effects [17]. Porphyrin has been developed as a smart material for drug delivery, which can potentially decrease the side effects of the target drug, increasing its efficient activities. Porphyrins are highly soluble in water and surfactants, and are frequently used in traditional photodynamic therapy. Several types of drugs of nanoscale design have been used together with porphyrins for a variety of applications in imaging and therapy [18,19]. This work was an extension to our previous study [1], to evaluate the Taxol yield by *Aspergillus flavipes* in response to different inhibitors, as well as to increase the bioactivity of extracted Taxol via chemical conjugation with porphyrin rings. In addition, the pharmacokinetic properties of native and modified Taxol were evaluated.

2. Results and Discussion

2.1. Taxol Production by the Selected Fungal Isolates in Response to Growth Inhibitors

Fungal endophytes elevated the hope for mass production of Taxol due to their fast growth, feasibility of fermentation, genetic manipulation, and independence of climatic changes [20]. Nevertheless, the anticipation of fungi for commercial Taxol production has been challenged by their lower yield and drastic loss of Taxol yield with sub-culturing and storage [5,6,8,13,21]. Thus, exploring novel fungal isolates with Taxol-producing potency was the objective. Four fungal isolates, namely, *Aspergillus flavipes* [1], *A. terreus*, *A. flavus*, and *A. parasiticus* [6], were selected based on their higher Taxol productivity. The isolates were grown on potato dextrose broth (PDB) and incubated, and then Taxol was extracted and quantified according to [1,6,8]. The yield of Taxol was emphasized by the TLC and HPLC analyses (Figure 1). The highest Taxol yield was assessed for *A. flavipes* cultures (185 µg/L), followed by *A. terreus* (66 µg/L), *A. flavus* (64 µg/L), and *A. parasiticus* (4 µg/L) (Figure 1). In an endeavor to increase Taxol yield by the selected fungal isolates, different growth inhibitors such as fluconazole and AgNO₃ were used. After the 5th day of incubation, the fungal cultures were amended with fluconazole and AgNO₃ (0, 2, 10, and 20 µg/mL final conc.), and then incubation was continued for 20 days before Taxol was extracted and quantified. From the results (Figure 2), fluconazole at 10 µg/mL displayed a strongly positive effect on Taxol yield, estimated as about 1.5-fold compared to control cultures. However, there was no detectable effect on Taxol yield by the tested fungal isolates in response to AgNO₃ compared to control cultures. The effect of fluconazole as a growth inhibitor on shifting the fungal secondary metabolites’ machinery toward terpenoids biosynthesis has been reported.
The modulation of Taxol yield by *A. flavipes* and *A. terreus* in response to fluconazole is reasonable, which could be due to inhibition of sterol synthesis as a competitive pathway with Taxol biosynthesis, directing the geranylgeranyl pyrophosphate pool to Taxol biosynthesis. Similar findings reported that AgNO₃ induced the production of Baccatin III and total taxanes [22], and that at higher fluconazole concentrations, the growth of *A. flavipes* and Taxol yield were significantly reduced, suggesting the inhibition of lanosterol demethylase, causing accumulation of toxic lanosterol intermediates [23].

Sterol and terpenoids are competing pathways for the initial precursor geranyl phosphate; so, the increase of Taxol yield by the experimented fungi with fluconazole incorporation might be explained by blocking the sterol synthesis, accumulating toxic lanosterol intermediates, inhibiting the squalene synthase, thus directing the flux of fungal intermediates towards synthesis of Taxol building blocks “taxadiene”. It has been reported that Taxol production could be induced by growing fungal cells under stress, similarly to other secondary metabolites [24]. To validate the concept of an inhibitory effect of fluconazole on sterol biosynthesis, the ergosterol contents of the fungal biomass were estimated under the same conditions. The cellular ergosterol concentration of *A. flavipes* was slightly reduced upon use of fluconazole; however, there was no detectable effect on ergosterol concentration in response to AgNO₃, suggesting the inhibitory effect of fluconazole on sterol biosynthesis as a competitive pathway with diterpenoid synthesis, which is consistent with our previous study [5–8].
A. terreus was increased about 2.3-fold compared to control cultures (without fluconazole). The (GGPP) substrate was determined by TLC, and visualized by iodine vapor. From the results, the intensity of residual GGPP spots were reduced gradually with the addition of fluconazole compared to controls, assuming a significant increase in TDS activity of A. flavipes upon fluconazole addition compared to controls. From the results (Figure 3), the activity of TDS of A. flavipes was noticeably increased upon addition of fluconazole in a concentration-dependent manner. At a higher concentration of fluconazole (20 µg/mL), the activity of TDS by A. flavipes and A. terreus was increased about 2.3-fold compared to control cultures (without fluconazole).

Taken together, based on TDS activity, Taxol yield, and ergosterol concentration, it could be deduced that fluconazole modulates the Taxol biosynthetic machinery by blocking the competitive sterol pathway, which is consistent with our previous results [1,5,6,25].
After conjugation, Taxol–porphyrin residues were purified from unbounded Taxol by silica gel column chromatography. The purple-colored Taxol–porphyrin conjugates were collected and dried, and the degree of modification was checked by TLC and UV analysis (Supplementary data). From the results, the Taxol–porphyrin conjugates were eluted at slightly different mobility rates, normalizing to native Taxol. Furthermore, as seen from the UV spectra, a reasonable shift on the absorption maxima of the native Taxol and Taxol–porphyrin conjugates occurred. The lower mobility rate of Taxol–porphyrin conjugates compared to native Taxol suggests the successfulness of the covalent chemical conjugation of Taxol–porphyrin.

2.3. Conjugation of Taxol with Porphyrin

The major clinical limitations for Taxol usage are its poor solubility in water and higher dose requirements to get the desired activity. Thus, chemical modification of Taxol and improving the formulation delivery systems are the main targets [26]. Construction of novel drug-therapy systems against cancer has become a research hotspot in biochemistry [27,28], and porphyrin-modified paclitaxel prodrugs are one of the practical technologies for such drug formulations. On the other hand, taking advantage of the fluorescent porphyrin part, the location of nanoparticles in cells could be explicit. Taxol was extracted and purified from A. flavipes, the location of nanoparticles in cells could be explicit. Taxol was extracted and purified from A. flavipes, and dried, and the degree of modification was checked by TLC and UV analysis (Supplementary data). From the results (Figure 4), the Taxol–porphyrin conjugates were eluted at slightly different mobility rates, normalizing to native Taxol. Furthermore, as seen from the UV spectra, a reasonable shift on the absorption maxima of the native Taxol and Taxol–porphyrin conjugates occurred. The lower mobility rate of Taxol–porphyrin conjugates compared to native Taxol suggests the successfulness of the covalent chemical conjugation of Taxol–porphyrin.
2.4. Antiproliferative Activity of Native Taxol and Taxol–Porphyrin Conjugates

The antiproliferative activity of the native *A. flavipes* Taxol and Taxol–porphyrin conjugates was evaluated against the liver carcinoma (HEPG2) and African green monkey kidney epithelial cell normal cells (VERO) using MTT assay, as described in Materials and Methods. The antiproliferative activity was expressed from the IC_{50} values (concentration of Taxol required to reduce the growth of 50% of initial number of tumor cells normalizing to control). From the results (Figure 5), the IC_{50} value of the native Taxol and Taxol–porphyrin conjugates was 60.7 µg/mL and 40.7 µg/mL, respectively. So, the antiproliferative activity of Taxol towards HepG2 tumor cells was increased about 1.5-fold upon conjugation with porphyrin. Interestingly, the toxicity of Taxol was reduced about 3-fold to normal VERO cells, as assessed from the IC_{50} values for native Taxol (113.9 µg/mL) and Taxol–porphyrin conjugates (312.2 µg/mL). Thus, it could be deduced that upon conjugation with porphyrin, the antiproliferative activity of Taxol against HepG2 cells was strongly increased; however, its cytotoxic activity was significantly reduced. This increase in anticancer activity with profound reduction to its cytotoxic effect could be ascribed to the molecular improvement on solubility of Taxol by increasing its surface hydrophilicity [29].

Figure 4. Conjugation of purified Taxol from *A. flavipes* with porphyrin. (A) Scheme of chemical conjugation of Taxol with porphyrin, as described in Materials and Methods. (B) TLC chromatogram of native Taxol and Taxol–porphyrin conjugates, (C) UV spectral analysis of native Taxol, Taxol–porphyrin conjugates, and unbounded porphyrin. The HPLC chromatogram of native Taxol (D) and Taxol–porphyrin conjugates (E).
were selected in the current study based on their yield of Taxol from our previous studies \[6,8\].

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The current fungal isolates were morphologically identified based on their macro and microscopical features \[30\].

2.5. Biochemical Profile of Mice in Response to Native and Taxol–Porphyrin Conjugates

The biochemical profile of mice in response to a single dose of native and Taxol–porphyrin conjugates was evaluated. From the data (Table 1), all of the measured biochemical parameters were slightly affected in response to Taxol–porphyrin conjugates, unlike to the significant changes due to unmodified Taxol, suggesting the mitigating and ameliorating effect of porphyrin on the Taxol compounds. A significant decrease in kidney enzymes, especially alanine amino transferase (ALT) and aspartate amino transferase (AST), in contrast to the significant decrease in alkaline phosphatase activity in response to unmodified Taxol was observed.

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<td>248</td>
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<td>Urea (mg/dl)</td>
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3. Materials and Methods

3.1. Fungal Isolates, Media Preparation, and Screening for Taxol Production

Four different fungal isolates, namely, Aspergillus flavipes, A. terreus, A. flavus, and A. parasiticus, were selected in the current study based on their yield of Taxol from our previous studies \[6,8\]. The current fungal isolates were morphologically identified based on their macro and microscopical features \[30\]. Aspergillus flavipes was isolated from Egyptian soil samples \[1,31–36\], while A. terreus, A. flavus, and A. parasiticus were isolated as endophytes from Podocarpus gracilior \[6,8\]. The isolates were maintained on potato dextrose agar (PDA) medium \[37\] (extract of 250 g potato, 20 g glucose, and 20 g agar, dissolved in 1 L tap water). The fungal isolates were screened for Taxol production by maintaining on potato dextrose agar (PDA) medium \[37\] (extract of 250 g potato, 20 g glucose, and 20 g agar, dissolved in 1 L tap water).
agar, dissolved in 1 L tap water). The fungal isolates were screened for Taxol production by inoculating 2 agar plugs of 5 mm from 6-day-old pure cultures of each fungal isolate into 50 mL/250 mL Erlenmeyer flask of potato dextrose broth (PDB) [38]. Three biological replicates from each fungal isolate were prepared. Flasks of PDB media without fungal inocula were used as negative controls at the same conditions. After incubation, mycelia were filtered from the liquid medium using sterile cheesecloth to remove the fungal mycelia, and fatty acids were precipitated from the filtrate by 0.03% sodium bicarbonate. Taxol was extracted with double volume of dichloromethane (DCM) and separated by TLC analysis as described previously [1–6].

3.2. Taxol Purification by Preparative-HPLC System

The extracted Taxol was purified by a semipreparative technique with Puriflash 4100 system (Interchim, Montluçon, France) with HPLC quaternary pump, a PDA–UV-Vis detector 190–840 nm, a fraction collector, and a sample loading module. For system controlling and process monitoring, Interchim software 5.0 (Montluçon, France) was used. The crude extract (3 g) was dissolved in 50 mL methanol, and then dissolved on a small amount of stationary phase (silica). The deposit solvent was evaporated with a rotary evaporator until dry powder was obtained. The fractionation was performed using automated flash chromatography (Puriflash 4100, Interchim), with pre-packed INTERCHIM PF-30SI-HP (Montluçon, France) (30 µm silica gel) columns at a flow rate of 15 mL/min, a pressure 22 bar, using a gradient of methanol and DCM for the elution.

3.3. TLC, UV, and HPLC Analyses

The sample was fractionated and identified by TLC [6,8,39] using Merck 1 mm (20 × 20 cm) pre-coated silica gel plates. The solvent systems chloroform/methanol (7:1, v/v), chloroform/ acetonitrile (7:3 v/v), ethylacetate/2-propanol (95:5 v/v), methylene chloride/tetrahydrofuran (6:2 v/v), and methylene chloride/methanol/dimethyl-formamide (90:9:1 v/v/v) were used. Taxol spots were detected by UV illumination (Min-UVIS, DESAGA, Heidelberg, Germany) at 254 nm regarding the Rf value of authentic Taxol (Cat. # T7402) giving blue-colored spots. The intensities of the target spots were assessed by the ImageJ software package (https://imagej.nih.gov/ij/download.html). The putative Taxol spots of silica were scraped off from the plate and eluted with DCM. The purity and concentration of eluted Taxol samples were analyzed by HPLC (Agilent Technol, G1315D, USA) with an RP- C18 column (Eclipse Plus C18 4.6 × 150 mm, 3.5 µm, # 959963-902). The isocratic mobile phase was acetonitrile/water (52:48 v/v/v) at a flow rate of 1.0 mL/min. The sample and mobile phase were filtered, injected into the column, and analyzed for 20 min. The fractions were scanned from 200 to 500 nm by a photodiode array detector (DAD). The identity and concentration of the Taxol sample were confirmed from the retention time and absorption peak area at 228 nm [21].

The UV spectra of recovered Taxol were scanned at λ 200–300 nm (Spectrophotometer, RIGOL, ultra-3000 Series, Hyperions ApS Sandholmvej, DK-3450 Allerød, Denmark) with blank medium as negative control. The samples were dissolved in methanol.

3.4. LC–MS Analyses

The spots of silica gel containing Taxol on the TLC sheets were scraped-off and eluted with methanol [6]. The chemical structure of extracted fungal Taxol was validated from LC–MS/MS analyses (Waters ACQUITY, LC-MS, XEVO-TQD#QCA423, Waters, USA). The HPLC portion was run isocratically with acetonitrile/water (49:51) as the mobile phase. The samples in 100% methanol were infused into the mass spectrometer through a reverse-phase C18 column at a flow rate of 0.2 mL/min with a column temperature of 27.8 ºC and a spray voltage of 3 kV by the loop injection method. The MS scanning ranged from 100 to 1000 m/z.
3.5. Eliciting the Taxol Yield of Selected Fungal Isolate with Growth Inhibitors

The effect of fluconazole on inducing the Taxol yield by the selected fungal isolate was evaluated [1]. The Taxol production media PDB was supplemented with different concentrations of fluconazole (0, 2, 10, and 20 µg/mL) after 5 days of preincubation. The cultures were re-incubated for 15 days, and then the filtered using sterile cheesecloth to remove the fungal mycelia. Taxol was extracted and assessed by TLC and HPLC, and the activity of Taxadiene synthase “rate-limiting enzyme” of Taxol biosynthesis was determined [8].

3.6. Taxadiene Synthase (TDS) Activity Assay

After incubation at the desired conditions, the fungal cultures were filtered and the collected mycelia were washed using sterile distilled water. Five grams of the fungal fresh biomass were pulverized in liquid nitrogen, dispensed in 10 mL of 30 mM, pH 8.0 HEPES buffer with 5 mM sodium ascorbate, 5 mM DTT, 5 mM Na₂S₂O₅, 15 mM MgCl₂, 10% (v/v) glycerol, and 1% (w/v) polyvinylpolypyrrolidone. The homogenate was centrifuged at 8000 rpm for 15 min; then, the supernatant was used as a source of crude enzyme. The activity of Taxadiene synthase (TDS) was assessed [40] with slight modifications [8]. The reaction mixture contained 500 µL of the crude enzyme, 1 mL of 30 mM HEPES buffer (pH 8.0), and 50 µL of 1 mM geranylgeranyl pyrophosphate (GGPP). The reaction was incubated at 32 °C for 1 h, and terminated by 50 µL of EDTA (0.5 M, pH 8.0). The GGPP concentration was determined by TLC using silica gel plates 60 F₂₅₄ (Merck KGaA, Darmstadt, Germany) with developing solvent system propan-2-ol, ammonia, and water (9:3:1) regarding to the authentic GGPP (Cat#. G6025) after visualization by vapor iodine [41]. The intensity of GGPP spots was determined by ImageJ software [1,6,8].

3.7. Total Sterol Assay

Total cellular sterols were extracted according to Breivik and Owades [42]. Briefly, 1.5 g homogenized fungal tissues was homogenized in alcoholic potassium hydroxide solution (4 mL), in 100 mm sterile borosilicate glass screw-cap tubes, with incubation at 85 °C for 1 h, and then the tubes were cooled to room temperature. Sterols were then extracted with n-heptane with vigorous mixing for 3 min. The heptane layer was transferred into a clean borosilicate glass screw-cap tube and stored at −20 °C until use. Prior to analysis, the sterol extract was diluted 5-fold in 100% ethanol and scanned at 240–300 nm (RIGOL, ultra-3000 Series, Hyperions ApS Sandholmvej, DK-3450 Allerød, Denmark). The presence of ergosterol and sterol intermediate 24(28) dehydro-ergosterol in the extract was confirmed from the distinctive four-peaked curve. A flat line indicated the absence of detectable ergosterol, and the amount of ergosterol was calculated as described previously [1,8].

3.8. Conjugation of Taxol with Porphyrin

The chemical conjugation of Taxol with porphyrin was conducted according to [17,29,43], as illustrated in Figure 4A. Briefly, tetracarboxyphenyl porphyrin (0.3 mM), N,N′-dicyclohexylcarbodi imide (DCC) (0.25 mM), 4-dimethylaminopyridine (DMAP) (0.1 mM), and Taxol (0.2 mM) were dissolved in tetrahydrofuran (THF) (10 mL), and the mixture was then stirred in the dark at room temperature under nitrogen conditions for 72 h. The solution was filtered, and the filtrate was dried under the reduced pressure to remove the solvent. The residues were dissolved in chloroform (50 mL) and washed with water (3 × 50 mL), and the organic phase was then dried over Na₂SO₄. The solvent was removed under the reduced pressure, and the crude product was purified by column chromatography to give Taxol–porphyrin conjugates as a purple solid pellet.

3.9. Antiproliferative Activity of Native Taxol and Taxol–Porphyrin Conjugates

The antiproliferative activity of native fungal Taxol and Taxol–porphyrin conjugates against liver carcinoma (HEPG2) and normal African green monkey kidney cell (VERO) was evaluated (VACSERA,
Cairo, Egypt). The tumor cell viability was assessed with 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide (MTT) assay [44]. The 96-well plate was seeded with 10^3 cells/well, incubated for 24 h, supplemented with different concentrations of native and Taxol–porphyrin conjugates, and re-incubated for 48 h. Twenty microliters (µL) of MTT dye was added, incubated for 6 h, and the developed purple color formazan complex was measured at λ570 nm. The IC_{50} value was expressed by the concentration of Taxol reducing the growth of 50% of initial number of cells, normalizing to positive control (Paclitaxel, No. 23958/2005, Cipla, India).

3.10. Biochemical Profile of Mice in Response to Native and Taxol–Porphyrin Conjugates

The biochemical toxicity of the native and Taxol–porphyrin conjugates was assessed in vivo using male mice (25 g, 30 days old). The experiments were conducted according to the guidelines of the Institutional Animal Care and Use Committee (IACUC) at Faculty of Medicine, Zagazig University, following the National Institute of health guidelines under protocol 15-08-263. After acclimatization for 3 days, a single dose (100 µL) of the native and Taxol–porphyrin conjugates (0.4 µg/mL) was injected into the tail vein of mice and acclimated for 20 days. Each treatment group had five mice. Normal mice and primate plasma of injected mice were used as negative and positive controls, respectively. Blood samples were collected intervally to assess the biochemical and hematological parameters. The cytotoxicity of Taxol and Taxol–porphyrin conjugates was determined based on biochemical parameters such as ALT, AST, ALP, total protein, albumin, globulin, urea, and creatinine (Spectrum Co., Cairo, Egypt).

3.11. Deposition of the Fungal Isolates

The potent Taxol-producing fungal isolates Aspergillus flavipes, A. terreus, and A. flavus were molecularly identified based on their sequences of 18S-ITS-28S regions and deposited in GenBank with accession numbers JF831014.1, MH156195, and MF377553, respectively.

3.12. Statistical Analyses

All experiments were conducted in triplicates and the data were analyzed using one-way ANOVA (https://goodcalculators.com/one-way-anova-calculator/) and represented by the means and standard deviations.

4. Conclusions

In conclusion, Taxol productivity by the four selected fungal isolates, namely, A. flavipes, A. terreus, A. flavus, and A. parasiticus, was evaluated in response to fungal growth inhibitors, fluconazole and silver nitrate. Maximum Taxol yield was detected with A. flavipes cultures, and its yield was duplicated upon addition of fluconazole. To increase the solubility of Taxol, it was conjugated with porphyrin, and the conjugation efficiency was evaluated by TLC and UV spectral analysis. The anticancer activity of Taxol was increased about 2-fold, upon conjugation with porphyrin, towards HepG2. Additionally, the cytotoxicity of Taxol–porphyrin conjugates on normal VERO cells was strongly reduced compared to native Taxol.

Supplementary Materials: The supplementary materials are available online.

Author Contributions: A.S.A.E.-S., M.F., B.S. conceived and designed the experiments, and wrote the manuscript; S.M., M.A.Y. and N.Z. performed the experiments; A.S.A.E.-S., M.F., M.S. and B.S. interpreted and revised the manuscript. All authors have read and agreed to the published version of the manuscript.

Funding: The authors appreciate the financial support from the Egyptian Academy of Scientific Research and Technology (ASRT-2017-20) (A.S.A.E.-S.), and County Council of Vasterbotten and Lions Cancer Research Fund (B.S.).

Conflicts of Interest: The authors declare no conflict of interest.
References


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**Sample Availability:** Samples of the Porphyrin-Taxol conjugates are available from the authors.