

Production of Major Mycotoxins by *Fusarium* Species Isolated from Wild Grasses in Peninsular Malaysia

(Penghasilan Mikotoksin Utama oleh Spesies *Fusarium* yang Dipencilkan daripada Rumput Liar di Semenanjung Malaysia)

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ABSTRACT

The *Fusarium* species are notoriously known for causing various plants and animal diseases and producing a number of harmful mycotoxins. The mycotoxins production by species recovered from non-agricultural hosts such as wild grasses have hitherto never been given attention. We examined 30 strains representing 12 *Fusarium* species i.e. *F. oxysporum*, *F. solani*, *F. semitectum*, *F. nelsonii*, *F. compactum*, *F. equiseti*, *F. chlamydosporum*, *F. proliferatum*, *F. subglutinans*, *F. sacchari*, *F. lateritium* and *F. incarnatum-equiseti* species complex isolated from wild grasses in Peninsular Malaysia for the production of four major mycotoxins i.e. moniliformin (MON), fumonisin B₁ (FB₁), zearalenone (ZEN) and beauvericin (BEA) using TLC and HPLC techniques. BEA was the highest frequency of mycotoxin detected, followed by MON, ZEN and FB₁. This study also presented the first report of BEA production by *F. solani*, *F. compactum* and *F. chlamydosporum*. All mycotoxins were not produced by *F. nelsonii* and *F. lateritium*. All *Fusarium* species were isolated from asymptomatic grasses, hence they are likely to exist as endophytes or latent pathogens.

Keywords: *Fusarium*; grasses; mycotoxins; Peninsular Malaysia

ABSTRAK

Spesies *Fusarium* dikenali sebagai penyebab pelbagai penyakit tumbuhan dan haiwan serta menghasilkan beberapa mikotoksin yang berbahaya. Penghasilan mikotoksin oleh spesies yang dipencilkan daripada perumah bukan pertanian seperti rumput liar, sehingga kini tidak pernah diberi perhatian. Kami memeriksa 30 strain mewakili 12 spesies *Fusarium* iaitu *F. oxysporum*, *F. solani*, *F. semitectum*, *F. nelsonii*, *F. compactum*, *F. equiseti*, *F. chlamydosporum*, *F. proliferatum*, *F. subglutinans*, *F. sacchari*, *F. lateritium* dan *F. incarnatum-equiseti* kompleks spesies yang dipencilkan daripada rumput liar di Semenanjung Malaysia untuk penghasilan empat mikotoksin utama iaitu moniliformin (MON), fumonisin B₁ (FB₁), zearalenon (ZEN) dan beauvericin (BEA) menggunakan teknik kromatografi lapisan nipis (TLC) dan kromatografi cecair berprestasi tinggi (HPLC). BEA merupakan mikotoksin yang paling kerap dikesan, diikuti oleh MON, ZEN dan FB₁. Kajian ini juga merupakan laporan pertama penghasilan BEA oleh *F. solani*, *F. compactum* dan *F. chlamydosporum*. Kesemua mikotoksin tidak dihasilkan oleh *F. nelsonii* dan *F. lateritium*. Semua spesies *Fusarium* tersebut dipencilkan daripada rumput yang tidak menunjukkan gejala penyakit, maka ia mungkin wujud sebagai endofit atau patogen pendam.

Kata kunci: *Fusarium*; mikotoksin; rumput; Semenanjung Malaysia

INTRODUCTION

Several species of the genus *Fusarium* are known for causing serious plant diseases on a number of economically important plants worldwide, including those in Malaysia such as corn (Darnetty et al. 2008), rice (Nur Ain Izzati et al. 2008), sugar cane (Siti Nordahliawate et al. 2008) and banana (Liew et al. 1998). Members of this genus also produce harmful secondary metabolites known as mycotoxins (Desjardins 2006) in food and feeds. The four most important mycotoxins produced by *Fusarium* are zearalenone (ZEN), moniliformin (MON), fumonisin B₁ (FB₁) and beauvericin (BEA) (Leslie et al. 2004; Logrieco et al. 2002; Sopterean & Puia 2012). The possible health risks on animals and humans have evoked global concern

over food safety and therefore, numerous research works have been focused on the toxicology of *Fusarium* mycotoxins (Lee et al. 2010; Negedu et al. 2011; Tan et al. 2012). ZEN has always been postulated to cause infertility among mammals, mammary hypertrophy in females and feminisation in males (Dacasto et al. 1995; Smith et al. 1994) with swine being the most vulnerable animal towards this toxin. Several studies showed MON is toxic, causing muscular weakness, respiratory distress, coma and even lead to fatality in tested animals (Engelhardt et al. 1989; Ledoux et al. 1995). The most recent study by Jonsson et al. (2012) and Sharma et al. (2012) have confirmed that heart is the main target tissue of MON toxicity in rats and avian. Among series of fumonisins, FB₁ is the most prominent

and frequently found in foods and feeds. Over the years, FB₁ has been linked to leukoencephalomalacia, a brain lesion in horses (Marasas et al. 1988), human oesophageal cancer (Sydenham et al. 1990), immunodepressive effects in turkey poults (Li et al. 2000) and several others diseases. Apart from the above mentioned mycotoxins, BEA is regarded as a new and less investigated secondary metabolites since its role as a toxin is poorly understood (Jestoi 2008). However, some studies showed that BEA has an insecticidal activity (Gupta et al. 1991), induces programmed cell death similar to apoptosis (Macchia et al. 2002) and causes chronotropic effect, a decrease in the frequency of cardiac spontaneous beating activity in guinea pig heart (Lemmens-Gruber et al. 2000).

In Malaysia, very limited number of studies on mycotoxin have been conducted. Desjardins et al. (1997) performed a study of FB₁ and MON production in rice infected by *G. fujikuroi* and the results showed the rice samples were contaminated with FB₁ and MON at concentration levels of 170 µg/g and 1000 – 5000 µg/g, respectively. Other studies were carried out by several researchers mostly on edible products e.g. on Malay traditional vegetables (Nur Ain Izzati & Wan Hasmida 2011), corns and animal feeds (Reddy & Salleh 2011) and cereals (Soleimany et al. 2011). All results showed certain amounts of mycotoxins contamination in tested samples. To date, there is no scientifically reliable data regarding mycotoxins production by *Fusarium* species from non-agricultural hosts, particularly wild grasses. Like any other plants, grasses are also suitable hosts for *Fusarium* species. Leslie et al. (2004) initiated a study on species diversity and mycotoxins production by *Gibberella fujikuroi* (*Fusarium* section Liseola) recovered from prairie grasses in Kansas and found generally low to moderate amounts of fumonisins (120 µg/g), BEA (4 – 320 µg/g) and fusaproliferin (50 – 540 µg/g). Nur Ain Izzati et al. (2009) have isolated and reported 10 *Fusarium* species i.e. *F. semitectum*, *F. solani*, *F. oxysporum*, *F. equiseti*, *F. sacchari*, *F. proliferatum*, *F. subglutinans*, *F. compactum*, *F. longipes* and *F. chlamydosporum* from 25 samples of wild grasses in Peninsular Malaysia; indicating the possibility for multiple mycotoxins production by these species *in planta*. Therefore, a well-defined attention should be given to the incidence, type and chemical natures of *Fusarium* toxins in this area since in Malaysia, wild grasses are commonly used as feeding stocks for ruminants which lead to health and safety issues.

Thus, the objective of this study was to examine the production of the four most common mycotoxins produced by *Fusarium* species isolated from wild grasses in Peninsular Malaysia, both qualitatively and quantitatively, with the aim of providing an additional data on mycotoxins occurrence in Malaysian samples.

MATERIALS AND METHODS

FUSARIUM STRAINS

Thirty strains of *Fusarium* isolated from 18 species of asymptomatic grasses (Table 1) collected throughout Peninsular Malaysia were examined for mycotoxins production *in vitro*. Strains were identified into species level following Leslie and Summerell (2006) as well as molecular approach described by O'Donnell et al. (1998), respectively. Strains were preserved in 15% glycerol (Salleh & Sulaiman 1984), catalogued and deposited at the *Fusarium* Culture Collection Unit, Universiti Sains Malaysia.

STANDARD SOLUTION OF MYCOTOXINS

All four mycotoxin standards were purchased from Sigma-Aldrich (St. Louis, MO, USA), diluted in methanol and prepared at concentrations ranging from 5 to 25 µg/g. The standard solutions were kept at 4°C for longer shelf life.

MYCOTOXINS PRODUCTION *IN VITRO*

Eighty five grams of corn grits (~ 45% moisture) were autoclaved in 250 mL Erlenmeyer flasks and inoculated with conidial suspensions (~1 × 10⁷ conidia/mL) of *Fusarium* strains grown on potato dextrose agar (PDA) for 7 days. Corn grits inoculated with sterilised distilled water served as controls. The corn grit cultures and controls were incubated at 25 ± 2°C for 28 days in total darkness.

MYCOTOXINS ANALYSES

Mycotoxins profiles of the 12 species of *Fusarium* strains were analyzed by using two techniques i.e. thin layer chromatography (TLC) for MON and FB₁ and high performance liquid chromatography (HPLC) for ZEN and BEA.

MONILIFORMIN (MON) AND FUMONISIN B₁ (FB₁)

The procedures of Burmeister et al. (1979) and Nelson et al. (1992) were adopted for extraction of MON and FB₁, respectively, with slight modifications. As for detection, plates were developed in a solvent system according to Thrane (1986) for MON and Nelson et al. (1992) for FB₁ and visualised under both white and UV (364 nm) lights. Retention factor (R_f) values for the standards and samples were calculated and compared.

ZEARALENONE (ZEN) AND BEAUVERICIN (BEA)

ZEN was extracted and analysed by techniques previously described by Bottalico et al. (1985) and Jimenez et al. (1997), with slight modifications. Meanwhile, BEA was extracted and analysed following the procedure of Logrieco et al. (1998) and Munkvold et al. (1998). The presence of

ZEN and BEA were confirmed and quantified by HPLC with a UV detector at 236 nm for ZEN and 205 nm for BEA at a flow rate of 0.6 mL/min. The extracts were injected into HPLC system and identified by comparing retention times and UV spectra of the samples with those of the standards and further quantified by comparing peak areas from the samples with a calibration curve of the standards.

RESULTS AND DISCUSSION

We examined 30 strains of selected *Fusarium* species recovered from 18 species of wild grasses in producing four major mycotoxins i.e. MON, FB₁, ZEN and BEA. Out of 30 strains of 12 *Fusarium* species recovered, nine strains of seven species produced detectable levels of MON i.e. *F. oxysporum* (T3507&N, M3548&N, P3610&N), *F. chlamydosporum* (P3590&N), *F. solani* (J3517&N), *F. proliferatum* (D3474&N), *F. subglutinans* (T3503&N), *F. sacchari* (T3671&N) and *F. incarnatum-equiseti* species complex (C3485&N) (Table 1). The extracts of the tested strains showed the same R_f value (~0.68) and colour (bluish) as MON standard and are in agreement with previous reports (Mubatanhema et al. 1999; Thrane 1986). All species have been reported as MON producer (Chelkowski et al. 1990; Lew et al. 1996). One strain represented *F. incarnatum-equiseti* species complex was able to produce MON. Further work is therefore required to confirm the ability of this species to produce this toxin as no previous data was reported on the production of MON by *F. incarnatum-equiseti* species complex. Meanwhile, MON was absent in cultures inoculated with five *Fusarium* species i.e. *F. semitectum*, *F. equiseti*, *F. compactum*, *F. nelsonii* and *F. lateritium*. Similar result was obtained by Jimenez et al. (1997) as they did not detect any trace levels of MON by *F. semitectum* in their study and claimed that this species was a low MON producer. Nur Ain Izzati and Wan Hasmida (2011) however managed to detect MON in corn grit cultures inoculated with *F. semitectum* isolated from traditional vegetables in Malaysia. There is no credible explanation has been made to clarify as to why some strains within the species are able to produce MON and some are not. Meanwhile, the four latter species are considered as non-producer of MON (Leslie & Summerell 2006).

FB₁ was only detected in one strain of *F. proliferatum* (strain D3474&N) and in agreement with previous study by Logrieco et al. (2002). The R_f value for FB₁ was 0.22 and the colour appeared as light purple under the white light and reddish spot under the long wave UV (364 nm). The fact that no FB₁ was detected in two *Fusarium* species in section Liseola i.e. *F. subglutinans* and *F. sacchari* was also a common phenomenon as both species have been reported to produce this mycotoxin at very low or undetectable levels in corn grit cultures (Leslie et al. 1996; Reynoso et al. 2004; Tseng et al. 1995).

ZEN was only produced in corn grits inoculated with two species i.e. *F. semitectum* (C3482&N) and *F. equiseti* (M3543&N). *F. semitectum* and *F. equiseti* have

been consistently classified as the main producers of ZEN (Frisvad & Thrane 2002; Hestbjerg et al. 2002). The concentration levels of ZEN produced by both species were low i.e. 2.8 and 4.4 µg/g, respectively and supported the proclamation by Kosiak et al. (2005) that countries with hot climate have not been presented with problems by ZEN contamination. Most strains of *F. oxysporum* did not produce ZEN; notwithstanding some reports accounted few strains of this species could produce ZEN (Marasas et al. 1984). The other 10 remaining species were reported as non-producers of ZEN (Desjardins 2006).

Nine species were able to produce detectable levels of BEA with low to moderate concentrations ranging from 19.5 to 567 µg/g. Two strains of *F. semitectum* (J3526&N and P3564&N), *F. equiseti* (D3741&N and C3494&N), *F. oxysporum* (T3507&N and P3610&N) and *F. proliferatum* (P3594&N and D3474&N) were positively detected for BEA and have been constantly reported as BEA producers (Leslie et al. 2004; Moretti et al. 2002). Meanwhile, only one strain each from *F. subglutinans* (T3503&N), *F. solani* (P3602&N), *F. compactum* (T3681&N) and *F. chlamydosporum* (D3696&N) produced BEA in the corn grit cultures. No earlier reports have been presented on the occurrence of BEA by *F. solani*, *F. compactum* and *F. chlamydosporum* (Leslie & Summerell 2006), as well as *F. incarnatum-equiseti* species complex. Hence, this study presented the first report of BEA production by these species. Several authors revealed that *F. subglutinans* was able to produce BEA in cultures (Leslie et al. 2004; Reynoso et al. 2004; Shephard et al. 1999) and in contrast with Moretti et al. (1996) and Munkvold et al. (1998) who found none BEA in *F. subglutinans* strains from corn samples in Iowa, Argentina and Italy. All strains of *F. sacchari* did not produce detectable levels of BEA. All four mycotoxins were apparently not produced by *F. nelsonii* and *F. lateritium*. Presumably the toxin profiles for this species is similar to those of the most closely related species from section *Arthrosporiella* i.e. *F. semitectum*. Meanwhile, *F. lateritium* was reported to produce other mycotoxins such as enniatins (Pieper et al. 1992) and lateropyrone (Bushnell et al. 1984).

CONCLUSION

Fusarium species isolated from wild grasses in Peninsular Malaysia were also able to produce the four major mycotoxins i.e. moniliformin (MON), fumonisin B₁ (FB₁), zearalenone (ZEN) and beauvericin (BEA). The results of this study may indicate a potential health risk for ruminants that feed on these grasses and consequently for humans who consume these animals as a protein source.

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TABLE 1. Production of four types of mycotoxins by 30 *Fusarium* strains isolated non-cultivated grasses in Peninsular Malaysia

<i>Fusarium</i> species ^a	Host ^b	Strains ^c	Location ^d	Mycotoxins (µg/g)
<i>F. semitectum</i>	<i>Chloris barbata</i>	C3482&N	Temerloh, Pahang	n.d
	<i>Paspalum conjugatum</i>	J3526&N	Skudai, Johor	n.d
	<i>Sporobolus diander</i>	P3564&N	Seberang Perai, P. Pinang	n.d
<i>F. solani</i>	<i>Digitaria setigera</i>	P3602&N	Teluk Bahang, P. Pinang	n.d
	<i>Axonopus compressus</i>	J3517&N	Skudai, Johor	+NA
	<i>Digitaria ciliaris</i>	D3477&N	Pasir Putih, Kelantan	n.d
	<i>Sporobolus diander</i>	D3741&N	Kubang Kerian, Kelantan	n.d
	<i>Eleusine indica</i>	M3543&N	Saint John, Melaka	+2.8
<i>F. compactum</i>	<i>Paspalum conjugatum</i>	C3494&N	Cameron Highlands, Pahang	n.d
	<i>Imperata cylindrica</i>	T3681&N	Kemaman, Terengganu	n.d
	<i>Cyanodon dactylon</i>	K3638&N	Padang Terap, Kedah	n.d
	<i>Axonopus compressus</i>	J3523&N	Skudai, Johor	n.d
	<i>Cenotheca lappacea</i>	T3507&N	Kemaman, Terengganu	+NA
<i>F. oxysporum</i>	<i>Eragrostis amabilis</i>	M3548&N	Saint John, Melaka	+NA
	<i>Lophatherum gracile</i>	P3610&N	Teluk Bahang, P. Pinang	+NA
	<i>Echinochloa crus-galli</i>	P3594&N	Relau, P. Pinang	n.d
<i>F. proliferatum</i>	<i>Eleusine indica</i>	M3542&N	Saint John, Melaka	n.d
	<i>Panicum repens</i>	D3474&N	Pasir Putih, Kelantan	+NA
	<i>Paspalum conjugatum</i>	T3514&N	Kemaman, Terengganu	n.d
<i>F. subglutinans</i>	<i>Paspalum orbiculare</i>	C3856&N	Cameron Highlands, Pahang	n.d
	<i>Cenotheca lappacea</i>	T3503&N	Kemaman, Terengganu	+NA
	<i>Eleusine indica</i>	K3619&N	Padang Terap, Kedah	n.d
<i>F. sacchari</i>	<i>Paspalum conjugatum</i>	J3527&N	Skudai, Johor	n.d
	<i>Chrysopogon aciculatus</i>	T3671&N	Kemaman, Terengganu	+NA
	<i>Eleusine indica</i>	D3696&N	Kuala Krai, Kelantan	n.d
<i>F. chlamydosporum</i>	<i>Pennisetum purpureum</i>	P3590&N	Relau, P. Pinang	+NA
	<i>Imperata cylindrica</i>	K3634&N	Padang Terap, Kedah	n.d
	<i>Echinochloa colona</i>	B3850&N	Sg. Besar, Selangor	n.d
<i>F. lateritium</i>	<i>Eragrostis amabilis</i>	M3550&N	Saint John, Melaka	n.d
	<i>Axonopus compressus</i>	C3485&N	Cameron Highlands, Pahang	+NA
	<i>F. incarnatum-equiseti</i> species complex ^e			n.d

^a*Fusarium* species identified according to Leslie and Summerell (2006); ^bHost of each *Fusarium* species isolated; ^cStrain, a coding system by *Fusarium* Collection Unit, Universiti Sains Malaysia, MALAYSIA; (initial alphabet = states in Malaysia; ampersand (&) = grasses/weeds, 'N' = non-pathogenic); ^dSampling sites in Peninsular Malaysia; ^eSpecies identified based on gene sequencing analyse; + = presence; n.d = not detected; NA = concentration values not available

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