Abstract. Background/Aim: Oxidative stress caused by the production of excessive cellular reactive oxygen species (ROS) and high levels of nitric oxide contribute to several human pathologies. This study aimed to examine the anti-oxidant effects of fusigen, a compound produced from Aureobasidium melanogenum. Materials and Methods: Extracts of A. melanogenum were selected as a source for the isolation of fusigen. The anti-oxidant, nitric oxide suppression, as well as the free radical scavenging activities of fusigen were tested in BEAS-2B human bronchial epithelial cell line (BEAS-2B cells) and human dermal papilla cells (DP cells) using specific fluorescence dyes and flow cytometry analysis. Cell viability was determined by the MTT assay. Results: Fusigen did not exert cytotoxicity in the human normal BEAS-2B and DP cells at concentrations up to 100 μM. Fusigen decreased basal levels of cellular ROS, as well as the levels of ROS induced by hydrogen peroxide and ferrous ion enrichment. ROS decreasing effect was confirmed in DP cells. In addition, fusigen treatment suppressed intracellular NO levels in both BEAS-2B and DP cells. Conclusion: The optimal process of production of purified fusigen from A. melanogenum was determined. Fusigen exhibited a low cytotoxic effect and the potential to suppress ROS and NO. These results demonstrated that fusigen may be used for the treatment or prevention of human diseases.

Reactive oxygen species (ROS) are the reactive derivatives of oxygen with or without radicals, such as hydrogen peroxide (H$_2$O$_2$), hydroxyl radical (•OH), and superoxide radical (O$_2$•–). These harmful ROS could be produced everywhere in our body and in all cells, and exert cytotoxicity by attacking necessary biomolecules including DNA, RNA, proteins and lipids (1). Normally, cells manage ROS through the production of antioxidants. However, the excess ROS could cause oxidative stress in cells, leading to cell death or disease (2). Hence, providing antioxidants to cells could be another way to control ROS.

Iron (II) or ferrous (Fe$^{2+}$) has single electron transfer competency that can cause many radical reactions (3). Furthermore, the excessive free iron mediates the production of highly reactive radicals, namely hydroxyl radicals, through the Fenton reaction (4, 5). The direct DNA damage caused by the interaction between hydroxyl radicals and DNA is well known to cause cell and tissue damage (6) leading to several human diseases. Therefore, iron chelators might be one of the solutions or key substances to control cellular hydroxyl radicals. Nitric oxide (NO) is a cellular mediator that has been involved in physiology and pathology. As NO is produced by the immune cells in order to destroy invading microorganisms, it may also be toxic to the neighboring cells and tissues (7).
In addition, NO can directly interact with free radicals to generate highly toxic molecules (8). Thus, researchers have reported on the promising role of NO inhibitor and NO scavengers in protecting against certain forms of tissue injury and inflammation (9-12).

Siderophores are natural small iron chelating ligands produced by many microorganisms including bacteria and fungi (13). Nowadays, more than 500 structures of siderophore have been reported in databases (14). Many siderophores have been studied regarding their medical applications, for instance, selective drug delivery, iron overload diseases treatment, haemochromatosis treatment, antimalaria activity, removal of trans element, as well as, free radical scavenging activity (15, 16). Hence, there is a possibility that siderophores act as antioxidants. Moreover, some hydroxamic siderophores can bind to nitric oxide (NO) and decrease mortality in a septic shock model. *Aureobasidium melanogenum* is a yeast-like fungus which can be found in diverse environments (17-21). This species can produce many valuable products, such as amino acids were studied for enhancing the siderophore production.

### Materials and Methods

**Materials.** The human bronchial epithelial BEAS-2B (BEAS) cells were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA). Immortalized dermal papilla cells (DP cells) were obtained from the American Type Culture Collection (Rockville, MD, USA).

**Methods.** Various combinations of carbon and nitrogen sources were used into iron deplete media to determine the best carbon and nitrogen sources for siderophore production. Optimum concentration of the best carbon and nitrogen sources was investigated using a central composite design ( CCD) and response surface methodology (RSM). Additional supplements such as amino acids were studied for enhancing the siderophore production.

**Siderophore production and extraction.** A loop full of *A. melanogenum* VK02 was inoculated into iron depleted medium at 25°C, under continuous stirring at 200 rpm for 2 days. The yeast cells (5x10⁷ cells) were transferred to 100 ml of siderophore production medium (7% sucrose, 1.1% (NH₄)₂SO₄, 0.3% K₂HPO₄, 0.1% citric acid, 0.008% MgSO₄, 0.0002% ZnSO₄, and 1.5 mM L-ornithine) and cultured for 5 days at 25°C, and 200 rpm. The culture was centrifuged at 13,000 rpm at 4°C for 20 min. The supernatant was reduced 10 folds on a rotary evaporator at 40°C. Three volumes of cold ethanol were added into the concentrated supernatant and vigorously shaken before allowing precipitation to occur in the freezer overnight. The mixture was filtrated to remove the precipitate following by solvent extraction according to the methods described by Neilands (13) and Wang (23) without FeCl₃ supplementation.

**Fusigen purification.** Extracted crude siderophore was concentrated to several milliliters before being loaded onto a Zetadex LH-20 packed column (3x150 cm). Water was used as a mobile phase. Siderophore in the fractions was detected using ferric perchlorate assay and pooled together before being concentrated to several fractions. Each fraction was reduced on a rotary evaporator at 40°C. Three volumes of cold ethanol were added into the concentrated supernatant and vigorously shaken before allowing precipitation to occur in the freezer overnight. The mixture was filtrated to remove the precipitate following by solvent extraction according to the methods described by Neilands (13) and Wang (23) without FeCl₃ supplementation.

**Screening of siderophore production.** A number of *Aureobasidium melanogenum* isolates from the Plant Biomass Utilization Research Unit (PBURU), Chulalongkorn University (Bangkok, Thailand) culture collection were screened for siderophore production. Each isolate of *A. melanogenum* was cultivated in iron depleted medium (2.5% sucrose, 0.4% (NH₄)₂SO₄, 0.3% K₂HPO₄, 0.1% citric acid, 0.008% MgSO₄, and 0.0002% ZnSO₄) at 25°C, under continuous stirring at 200 rpm for 5 days. Cultured medium was collected and cells were discarded by centrifugation. The siderophore concentration was measured using a ferric perchlorate assay and monitoring absorbance at 450 nm.

**ROS detection.** The accumulated intracellular ROS were assessed using the fluorescent probe DCFH-DA and flow cytometry. In brief, cells were trypsinized and washed before being resuspended and incubated with 10 μM DCFH-DA and different concentrations of fusigen in serum free DMEM for 30 min at 4°C. DCFH-DA incubated cells were treated with 100 μM H₂O₂ and 150 μM FeSO₄ for 1 h at 37°C. Cells were washed, resuspended and immediately measured for green fluorescence intensity by Guava® easyCyte flow cytometer (Merck,
Nitric oxide detection. The intracellular NO was analyzed by flow cytometry using DAF-FM DA as a NO-specific fluorescent probe. Cells were trypsinized and then incubated with 10 μM DAF-FM DA in serum free DMEM for 30 min at 4°C, and subsequently at 37°C for 2 h. After incubation, cells were trypsinized, washed, resuspended, and immediately measured for green fluorescence intensity by Guava® easyCyte flow cytometer (Merck, Darmstadt, Germany). Mean of fluorescence intensity was quantified by Guava® InCyte software (Merck) analysis of the record histogram.

Statistical analysis. Data were expressed as the means±SD or SE from three or more independent experiments. Statistical analysis was performed by ANOVA at a significance level of p<0.05.

Results

Fusigen production from A. melanogenum. After screening 69 isolates of A. melanogenum from the PBURU culture collection of siderophore production, only A. melanogenum VK02 isolate from greasy aluminum surface in Ratchaburi province of Thailand could produce a superior yield of siderophore. The best siderophore production conditions and the process that could enhance siderophore production is shown in “Materials and Methods” section and Figure 2. The siderophore fusigen (chemical structure shown in Figure 1) was purified by flowing through Zetadex LH-20 column and C\textsubscript{18} reverse-phase column. The purified fusigen was analyzed by C\textsubscript{18} reverse-phase HPLC as described in “Materials and Methods” section. The purity of the fusigen is shown by the HPLC analysis, which shows only one clear peak at retention time of 10.063 min (Figure 3).

Effect of fusigen on viability of cells. To determine toxicity of the purified fusigen (Figure 2), human normal epithelial BEAS and human DP cells were incubated with various concentrations of fusigen. After 24 h, cell viability was determined by MTT assay. Figure 4 shows that fusigen caused no cytotoxicity to BEAS and DP cells at concentrations up to 100 μM. In order to test the possible effects of the compound on the levels of intracellular ROS and NO, fusigen was used at the concentrations of 0, 50, and 100 μM.

Fusigen inhibition of accumulated ROS generation. BEAS and DP cells were exposure to the DCFH-DA fluorescence dye for 30 min in the presence or absence of 50-100 μM fusigen and intracellular ROS levels were then analyzed by flow cytometry after 1 h. Figure 5 shows that the DCFH-DA fluorescent intensities in fusigen (50 and 100 μM) treated cells were significantly lower than those of control non-treated cells. In addition, fusigen showed strong anti-oxidant activity against high levels of oxidative stress induced by the treatment of BEAS cells with hydrogen peroxide (100 μM) and FeSO\textsubscript{4} (150 μM) plus hydrogen peroxide (100 μM). The antioxidant effect of the compound was confirmed in the DP cells treated with hydrogen peroxide.
Effect of fusigen on NO levels. Like ROS, NO has been shown to contribute to several human pathologies. Therefore, the effect of fusigen on NO production in epithelial and DP cells was examined. Flow cytometric evaluation of intracellular NO levels was performed using DAF-FM DA as a NO specific probe. The cells were incubated with the fluorescence probe for 30 min and the intracellular NO level was determined after 2 h. Figure 6 shows that the levels of NO in BEAS and DP cells were spontaneously up-regulated and the treatment of the cells with fusigen at the concentrations of 50 and 100 μM attenuated the increase of NO.

Discussion

Oxidative stress and reactive nitrogen species have been recognized as important accelerators of human diseases. In particular, the oxygen-derived radicals like hydroxyl radical and others have been shown to cause DNA damage that may contribute to cancers (24, 25). In addition, neurodegenerative diseases as well as the inflammatory related diseases have been shown to be linked with excessive production of cellular ROS (26-28). They are involved in production and aggregation of Aβ in Alzheimer’s disease (29). Moreover, they play a role in Parkinson’s disease by inducing apoptosis. In cancer, ROS play a role in DNA damage, initiation of tumorigenicity, and enhancement of invasion and metastasis (30-32). These studies have highlighted the importance of potent anti-oxidant compounds for the treatment or prevention of ROS-related diseases.

Here, we invented the process of fusigen production. The effect of culture medium on siderophore production from A. melanogenum VK02 was studied using various carbon sources including glucose, fructose, xylose, sucrose, and soluble starch, and nitrogen sources including (NH₄)₂SO₄, NH₄NO₃, NaNO₃, KNO₃, and urea. Various concentrations of L-glutamate, L-ornithine, and L-arginine were also used. The best conditions for siderophore production are found. Sucrose and (NH₄)₂SO₄ at 7% and 1.1% (w/v), respectively, were found to be the best carbon and nitrogen sources for siderophore production. Three amino acids were selected as the precursors for siderophore biosynthetic pathway. Only L-ornithine which is the key precursor of siderophore synthesis, enhanced siderophore production in this isolate. This carbon source and amino acid were also the best for siderophore production from A. melanogenum HN6.2 isolate (33). In contrast, the type of nitrogen source was different. A. melanogenem HN6.2 isolate preferred ammonium nitrate as a nitrogen source, while VK02 isolate preferred ammonium sulfate. The siderophore produced from A. melanogenum has been reported earlier to be fusigen, which

Figure 2. Overall scheme of fusigen production.
is one of the hydroxamate-type siderophores (23). Fusigen is also known as fusarinine C. Since the initial study on siderophores, it has been reported to be a stereotype from *Fusarium* species (34, 35). The purity of fusigen produced from VK02 isolate was determined by reverse-phase HPLC. A single sharp peak at 10.063 min of retention time was obtained, showing no contamination with other compounds.

Our results showed that fusigen was not toxic to human normal cells (Figure 4) at concentrations up to 100 μM. At the non-cytotoxic concentrations, fusigen could inhibit basal ROS production and H$_2$O$_2$-induced ROS in BEAS and DP cells. Via Fenton reaction, hydrogen peroxide could be converted to highly reactive molecule hydroxyl radical. Hydroxyl radical is the most biologically active free radical formed under hypoxic conditions. Its formation could be promoted by iron and can be the cause of atherosclerosis (36, 37). Moreover, hydroxyl radical can induce polymerization of fibrinogen resulting in an insoluble fibrin-like precipitate. This precipitate is reported to be involved in some degenerative diseases such as atherosclerosis, cancer, and
Figure 5. ROS scavenging activity of fusigen in BEAS-2B and DP cells. Cells were treated with 10 μM DCFH-DA and various concentrations of fusigen (0, 50, and 100 μM) for 30 min at 4°C before treatment with ROS generators (100 μM H₂O₂ or 100 μM H₂O₂ +150 μM FeSO₄) for 1 h. ROS production was determined by flow cytometry. Columns are mean±SE (count event=5,000). *Significantly different at p<0.05 compared to the untreated control group.
neurological disorders (38-41). The increase in cellular oxidative stress in the BEAS cells treated with hydroxyl radical generator could be attenuated by treating with fusigen, suggesting the use of this agent in the protection from harmful hydroxyl radicals. In addition, the NO suppressive effect of fusigen was also demonstrated (Figure 6). As NO may contribute to several human diseases including cancers and inflammation, this NO suppressive activity of fusigen may be useful for treating these conditions.

In conclusion, a novel process of fusigen production was presented as well as its activity in reducing cellular ROS and NO. Fusigen shows promising effects with low toxicity to normal cells. These findings support the use of fusigen in the treatment of oxidative stress-related diseases.

Conflicts of Interest

The Authors have no conflicts of interest to declare regarding this study.

Authors’ Contributions

Conceptualization, P.C.; Methodology, V.K., S.P., and P.C. Investigation, V.K., P.L., C.K., P.P., H.P., S.P., and P.C. Writing—original draft preparation, V.K. and P.C.; Writing—review and editing, P.C.; Funding acquisition, P.C.; Supervision, S.P. and P.C.

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References

1. Farber JL: Mechanisms of cell injury by activated oxygen species. Environ Health Perspect 102: 17-24, 1994. PMID: 7705293, DOI: 10.1289/ehp.94102s1017
2. Halliwell B and Gutteridge JM: Free radicals in biology and medicine. Oxford University Press, 2015. DOI:10.1093/acprof:oso/9780198717478.001.0001
3. AbouL-Enein AM, El-Baz FK, El-Baroty GS, Youssef A and Abd El-Baky HH: Antioxidant activity of algal extracts on lipid peroxidation. J Med Sci 3(1): 87-98, 2003. DOI: 10.3923/jms.2003.87.98
4. Halliwell B and Gutteridge JM: Free radicals and catalytic metal ions in human disease: An overview. Methods Enzymol 186: 1-85, 1990. PMID: 2172697, DOI: 10.1016/0076-6879(90)86093-B
5. Winterbourn CC: Toxicity of iron and hydrogen peroxide: The fenton reaction. Toxicol Lett 82-83: 969-974, 1995. PMID: 8597169, DOI: 10.1016/0378-4274(95)03532-X
6. Dzidzaroglu M and Jaruga P: Mechanisms of free radical-induced damage to DNA. Free Radic Res 46(4): 382-419, 2012. PMID: 22276778, DOI: 10.3109/10715762.2011.653969
7. Vallance P and Charles I: Nitric oxide as an antimicrobial agent: Does no always mean no? Gut 42(3): 313-314, 1998. PMID: 9577329, DOI: 10.1136/gut.42.3.313
8. Martinez-Ruiz A, Cadenas S and Lamas S: Nitric oxide signaling: Classical, less classical, and nonclassical mechanisms. Free Radic Biol Med 51(1): 17-29, 2011. PMID: 21549190, DOI: 10.1016/j.freeradbiomed.2011.04.010
Cauwels A: Nitric oxide in shock. Kidney Int 72(5): 557-565, 2007. PMID: 17538569, DOI: 10.1038/sj.ki.5002340

Lorente JA, Landin L, De Pablo R, Renes E and Liste D: L-arginine pathway in the sepsis syndrome. Crit Care Med 21(9): 1287-1295, 1993. PMID: 8370291

Petros A, Lamb G, Leone A, Moncada S, Bennett D and Vallance P: Effects of a nitric oxide synthase inhibitor in humans with septic shock. Cardiovasc Res 28(1): 34-39, 1994. PMID: 7509259, DOI: 10.1093/cvr/28.1.34

Pekidyshova DA, Bondarenko NA, Malyshchik I, Mikoian VD, Petros A, Lamb G, Leone A, Moncada S, Bennett D and Lorente JA, Landin L, De Pablo R, Renes E and Liste D: L-arginine pathway in the sepsis syndrome. Crit Care Med 21(9): 1287-1295, 1993. PMID: 8370291

Prasongsuk S, Sullivan R, Kuhirun M, Eveleigh D and Punnapayak H: Antioxidant and free radical scavenging activity of marine-derived strains of Aureobasidium pullulans. Folia Microbiol Res 3(5): 253-257, 2009.

Pokidyshev DA, Bondarenko NA, Malyshchik I, Mikoian VD, Petros A, Lamb G, Leone A, Moncada S, Bennett D and Lorente JA, Landin L, De Pablo R, Renes E and Liste D: The current status of siderophore production by the marine-derived Aureobasidium pullulans and its antimicrobial activity. Bioresour Technol 100(9): 2639-2641, 2009. PMID: 19162476, DOI: 10.1016/j.biortech.2008.12.010

Reuter S, Gupta SC, Chaturvedi MM and Aggarwal BB: Oxidative stress, inflammation, and cancer: How are they linked? Free Radic Biol Med 49(11): 1603-1616, 2010. PMID: 20840865, DOI: 10.1016/j.freeradbiomed.2010.09.006

Wang W, Chi Z, Chi Z, Li J and Wang X: Siderophore production by the marine-derived Aureobasidium pullulans and its antimicrobial activity. J Ren Nutr 21(3): 213-218, 2016. DOI: 10.1111/j.1432-1033.1967.tb19518.x

Pokidyshev DA, Bondarenko NA, Malyshchik I, Mikoian VD, Petros A, Lamb G, Leone A, Moncada S, Bennett D and Lorente JA, Landin L, De Pablo R, Renes E and Liste D: The current status of siderophore production by the marine-derived Aureobasidium pullulans and its antimicrobial activity. Bioresour Technol 100(9): 2639-2641, 2009. PMID: 19162476, DOI: 10.1016/j.biortech.2008.12.010

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