The cytotoxic effects of fumonisin B₁ in human kidney cells and
the ability of allicin to ameliorate these effects

By

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School of Laboratory Medicine and Medical Sciences

College of Health Sciences

University of KwaZulu-Natal, Durban

2019
DECLARATION

I Miss NNF Mahlalela declare that:

1. This dissertation contains original work done by the author and has not been submitted to UKZN or any other tertiary institution for the purposes of obtaining an academic qualification, whether by myself or any other party. The use of work by others has been duly acknowledged in the text.

2. The research described in this study was carried out in the Department of Medical Biochemistry and Chemical Pathology, School of Laboratory Medicine and Medical Science, Faculty of Health Sciences, University of KwaZulu-Natal, Durban, under the supervision of Dr R.B Khan.

Signed: ___________________________  Date: 19 July 2019
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<tr>
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<tr>
<td>AGE</td>
<td>Aged garlic extract</td>
</tr>
<tr>
<td>Apaf1</td>
<td>Apoptotic-protease-activating-factor-1</td>
</tr>
<tr>
<td>iROS</td>
<td>Intercellular reactive oxygen species</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>Bax</td>
<td>Bcl-2-associated X</td>
</tr>
<tr>
<td>BCA</td>
<td>Bicinchoninic acid</td>
</tr>
<tr>
<td>Bcl-2</td>
<td>B-cell lymphoma/leukemia-2</td>
</tr>
<tr>
<td>BH3</td>
<td>Bcl-2 homology domain 3</td>
</tr>
<tr>
<td>BHT</td>
<td>Butylated hydroxytoluene</td>
</tr>
<tr>
<td>BSA</td>
<td>Bovine serum albumin</td>
</tr>
<tr>
<td>CCM</td>
<td>Complete culture medium</td>
</tr>
<tr>
<td>cDNA</td>
<td>Complimentary DNA</td>
</tr>
<tr>
<td>CO₂</td>
<td>Carbon dioxide</td>
</tr>
<tr>
<td>CuSO₄</td>
<td>Copper sulphate</td>
</tr>
<tr>
<td>DADS</td>
<td>Diallyl disulfide</td>
</tr>
<tr>
<td>DAS</td>
<td>Diallyl sulfide</td>
</tr>
<tr>
<td>DAT</td>
<td>Diallyl trisulfide</td>
</tr>
<tr>
<td>DD</td>
<td>Death domain</td>
</tr>
<tr>
<td>dH₂O</td>
<td>De-ionised water</td>
</tr>
<tr>
<td>DISC</td>
<td>Death inducing signalling complex</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s Modified Eagle’s Medium</td>
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<tr>
<td>Term</td>
<td>Full Form</td>
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<td>------------------------------------</td>
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<tr>
<td>DMSO</td>
<td>Dimethyl sulphoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxymethylribose nucleic acid</td>
</tr>
<tr>
<td>DNase</td>
<td>Deoxyribonuclease</td>
</tr>
<tr>
<td>DNMT</td>
<td>DNA methyltransferase</td>
</tr>
<tr>
<td>EDTA</td>
<td>Ethylenediaminetetraacetic acid</td>
</tr>
<tr>
<td>Ex</td>
<td>Excitation wavelength</td>
</tr>
<tr>
<td>Em</td>
<td>Emission wavelength</td>
</tr>
<tr>
<td>FB1</td>
<td>Fumonisin B1</td>
</tr>
<tr>
<td>GPx</td>
<td>Gluathione peroxidase</td>
</tr>
<tr>
<td>GSH</td>
<td>Glutathione</td>
</tr>
<tr>
<td>H2O</td>
<td>Water</td>
</tr>
<tr>
<td>H2O2</td>
<td>Hydrogen peroxide</td>
</tr>
<tr>
<td>H3PO4</td>
<td>Phosphoric acid</td>
</tr>
<tr>
<td>HCl</td>
<td>Hydrochloric acid</td>
</tr>
<tr>
<td>HEK293</td>
<td>Human embryonic kidney cells</td>
</tr>
<tr>
<td>IARC</td>
<td>International agency for research on cancer</td>
</tr>
<tr>
<td>IC50</td>
<td>Median Inhibition Concentration</td>
</tr>
<tr>
<td>iCAD</td>
<td>Inhibitors of caspase DNase</td>
</tr>
<tr>
<td>LDH</td>
<td>Lactate dehydrogenase</td>
</tr>
<tr>
<td>LEM</td>
<td>Leukoencephalomalacia</td>
</tr>
<tr>
<td>LLC-PK1</td>
<td>Pig kidney epithelial cells</td>
</tr>
<tr>
<td>LMPA</td>
<td>Low melting point agarose</td>
</tr>
<tr>
<td>MDA</td>
<td>Malondialdehyde</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Name</td>
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<tr>
<td>--------------</td>
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</tr>
<tr>
<td>miRNA</td>
<td>MicroRNA</td>
</tr>
<tr>
<td>MTT</td>
<td>Methyl tetrazolium</td>
</tr>
<tr>
<td>NaCl</td>
<td>Sodium chloride</td>
</tr>
<tr>
<td>NaOH</td>
<td>Sodium hydroxide</td>
</tr>
<tr>
<td>NEDD</td>
<td>N-(1-naphthyl)ethylenediamine</td>
</tr>
<tr>
<td>Nm</td>
<td>Nanometer</td>
</tr>
<tr>
<td>Nrf2</td>
<td>Nuclear factor erythroid 2 (NFE2)-related factor 2</td>
</tr>
<tr>
<td>NTD</td>
<td>Neural tube defect</td>
</tr>
<tr>
<td>O₂⁻</td>
<td>Superoxide ion</td>
</tr>
<tr>
<td>OC</td>
<td>Oesophageal cancer</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate-buffered saline</td>
</tr>
<tr>
<td>PCR</td>
<td>Polymerase chain reaction</td>
</tr>
<tr>
<td>PFA</td>
<td>Paraformaldehyde</td>
</tr>
<tr>
<td>Prxs</td>
<td>Peroxidase</td>
</tr>
<tr>
<td>qPCR</td>
<td>Quantitative polymerase chain reaction</td>
</tr>
<tr>
<td>RLU</td>
<td>Relative light unit</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribonucleic acid</td>
</tr>
<tr>
<td>RNA</td>
<td>Ribose nucleic acid</td>
</tr>
<tr>
<td>RNS</td>
<td>Reactive nitrogen species</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>RT</td>
<td>Room temperature</td>
</tr>
<tr>
<td>SAC</td>
<td>S-allylcysteine</td>
</tr>
<tr>
<td>SAM</td>
<td>S-adenosyl methionine</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>SCGE</td>
<td>Single cell gel electrophoresis</td>
</tr>
<tr>
<td>SDH</td>
<td>Succinate dehydrogenase</td>
</tr>
<tr>
<td>SDS-PAGE</td>
<td>Sodium dodecyl sulphate - polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>SOD</td>
<td>Superoxide dismutase</td>
</tr>
<tr>
<td>SULF</td>
<td>Sulfanilamide</td>
</tr>
<tr>
<td>TBA</td>
<td>Thiobarbituric Acid</td>
</tr>
<tr>
<td>TBARS</td>
<td>Thiobarbituric Acid Reactive Substances</td>
</tr>
<tr>
<td>tBID</td>
<td>Truncated BH3 interacting-domain death agonist</td>
</tr>
<tr>
<td>TCA</td>
<td>Tricarboxylic acid</td>
</tr>
<tr>
<td>TNF</td>
<td>Tumor necrosis factor</td>
</tr>
<tr>
<td>TTBS</td>
<td>Tris-buffered saline containing 0.5% Tween20</td>
</tr>
<tr>
<td>VCl₃</td>
<td>Vanadium (III) chloride</td>
</tr>
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ABSTRACT

Fumonisin B₁ (FB₁) is a widespread contaminant of crops and is produced as a secondary metabolite of fungi. It has been found to disrupt sphingolipid metabolism, cause epigenetic modifications and induce cellular toxicity that can manifest through oxidative stress and apoptosis. Although implicated in animal toxicity including kidney cancer in rats, FB₁ effects in the human kidney have not been explored. Allicin is a biological component of garlic that has been widely studied for its health benefits including its anti-cancer and antioxidant properties. This study evaluated allicin as a possible therapeutic measure for FB₁-induced cytotoxicity in Hek293 cells. Both FB₁ and allicin decreased cellular viability (MTT assay) in a dose-dependent manner and generated IC₅₀ values of 215 µM and 3.905 µM respectively. Three 24 h treatments (FB₁, allicin and combined FB₁+allicin) were compared to untreated cells for induction of apoptosis and oxidative stress. Luminometry was used to determine cytotoxicity (lactate dehydrogenase assay), caspase activity and mitochondrial toxicity (apoptosis), and quantify intracellular ROS (iROS) and glutathione (GSH) (oxidative stress). Free radical production was estimated by the TBARS and NOS assays respectively, while DNA damage was evaluated using the comet assay. Western blotting confirmed the expression of various antioxidant and apoptotic proteins, and superoxide dismutase 2 (SOD2) transcripts were quantified using qPCR. ATP concentration was increased for all treatments, but mitochondrial toxicity was increased in the allicin treatment. While lipid peroxidation decreased, luminometry results indicate that iROS was increased and was accompanied by a corresponding decrease in SOD2 and catalase protein expression. Depletion of GSH was consistent with increased GPx1, HSP70 and Nrf2 protein expression suggesting the presence
of oxidative stress. SOD2 transcripts were only increased in the allicin treatment. Apoptosis was initiated as indicated by increased caspases 8 and 9, and pro-apoptotic Bax protein expression, but caspase 3/7 was not activated for the FB₁ treatment. However, DNA fragmentation and cPARP were increased for all treatments suggesting that apoptosis was executed. Overall, FB₁ and allicin individually and combined induced oxidative stress by increasing ROS and decreasing antioxidants. Apoptosis was also induced, although in a caspase independent manner in the FB₁ treatment. Overall, allicin did not ameliorate the effects of FB₁ in Hek293 cells.
CHAPTER 1: INTRODUCTION

1.1 Background

Plants form a vital part in the provision of energy to sustain normal metabolic function. However, the plant based foods consumed may contain harmful substances that arise due to contamination of the plant. Notable examples are toxic fungal metabolites such as fumonisin that are not removed by washing, heating or food preparation methods; exposure results in diverse toxicity ranging from immunotoxicity to cancer. Conversely, the organosulphur-containing herbal constituents that are added to enhance the flavour of food during cooking may confer health-benefits when consumed. Onion and garlic are among the plants that contain these useful substances. Garlic is renowned for its potential anti-cancer properties.

Fumonisin B (FB) mycotoxins are produced as secondary metabolites by fungi including *Fusarium verticillioides* (formerly called *Fusarium moniliforme*), *Fusarium proliferatum* and *Aspergillus niger*. They are known to contaminate crops such as maize, wheat and other grain products. Fumonisin B$_1$ (FB$_1$) is the most toxic member of the 28 types of the fumonisin family that have been characterised (Miller et al., 1993, Berek et al., 2001, Schmale and Munkvold, 2009, De Baere et al., 2018).

Human and animal exposure to FB$_1$ occurs via consumption of contaminated maize and other maize products. Exposure to FB$_1$ produces varying clinical symptoms from species to species and includes pulmonary oedema in pigs, leukoencephalomalacia in horses and oesophageal cancer (OC) in humans (Harrison et al., 1990, Kellerman et al., 1990, Van der Westhuizen et al., 2010). *In vivo* studies suggest the existence of fumonisin-induced carcinogenicity, immunosuppression, hepatotoxicity and nephrotoxicity in laboratory animals (Howard et al., 2001, De Baere et al., 2018).
When tested, FB₁ is neither genotoxic nor mutagenic according to standard protocols for genotoxic carcinogens, meaning it has no direct mechanism by which it affects DNA (Fink-Gremmels, 1999). The accepted mechanism for FB₁ cytotoxic effects is dysregulation of sphingolipid metabolism, however a study by Chuturgoon et al. (2014) has proposed an alternative mechanism of cytotoxicity which involves histone demethylation and DNA hypomethylation (Chuturgoon et al., 2014). Furthermore, FB₁ exposure has been shown to induce oxidative stress and thus play a role in carcinogenicity and toxicity. FB₁ was also found to activate apoptosis in many cell lines, primary cell cultures as well as in vivo (Domijan, 2012). An epigenetic mechanism of CpG promoter DNA methylation may play a big part in FB₁ toxicity as suggested by a study done on rat kidney and liver cells (Klarić et al., 2007, Sancak and Ozden, 2015, Demirel et al., 2015).

FB₁ is poorly absorbed and rapidly eliminated, a characteristic that could be attributed to the polar nature of this mycotoxin. Excretion occurs in both faeces and urine. In fact, urinary FB₁ may be used to assess human exposure to FB₁ and suggests ongoing exposure of the urinary system, including the kidney to FB₁. A study by Howard et al. (2001) using rats and mice found that FB₁ was a renal carcinogen in male rats’ diet at higher dosages (Howard et al., 2001), this finding supported a study that hypothesised that FB₁ can be carcinogenic at increased levels of exposure (Dragan et al., 2001). The inhibition of sphingolipid synthesis was accompanied by decreased cell proliferation in pig kidney epithelial cells (LLC-PK₁, renal proximal tubule epithelial cells developed from pigs) following exposure to fumonisins (Abado-Becogne et al., 1998). FB₁ was also found to target the proximal tubular epithelium cells in kidneys of rabbits and presented with toxin-induced renal failure (Gumprecht et al., 1995). It is therefore necessary to assess the effect of FB₁ in human kidney cells.

Garlic (Allium sativum) is a well-known food additive with many postulated health benefits that are attributed to the compound allicin (diallyl thiosulphinate) (Amagase, 2006). However, allicin is not present in garlic cloves; rather this unstable organosulphur compound is liberated when whole garlic
is macerated. Chopping, crushing or chewing garlic results in the release of alliinase, the enzyme responsible for the conversion of alliin to allicin. Allicin is found to have numerous biological effects such as anti-cancer, anti-microbial, anti-viral and anti-bacterial activities. Studies have shown that the anti-cancer properties may be due to inhibition of cancer cell proliferation or induction of apoptosis. In addition, allicin possesses powerful antioxidant properties (Hirsch et al., 2000, Oommen et al., 2004).

1.2 **Problem statement**
Consumption of fungal-contaminated maize, wheat and other grain products has adverse effects in animals and humans. The mycotoxin FB\textsubscript{1} predisposes different species to varying clinical symptoms including OC in humans and kidney cancer in rats. The study done by van der Westhuizen et al. (2010) on the prevalence of human OC in Southern Africa due to the exposure to FB\textsubscript{1} necessitates the study on the implication of FB\textsubscript{1} in kidney toxicity (Van der Westhuizen et al., 2010). While FB\textsubscript{1} dysregulates apoptosis and induces oxidative stress, allicin induces apoptosis and acts as an antioxidant. Therefore, the potential antagonistic effect of allicin as an inexpensive and implementable preventative measure against FB\textsubscript{1}-induced cytotoxicity warrants investigation.

1.3 **Hypothesis**
FB\textsubscript{1} induced cytotoxicity in human kidney cells may be reduced by co-exposure to allicin.

1.4 **Aim**
This study aimed to determine the cytotoxic effects of FB\textsubscript{1} in human kidney cells and the ability of allicin to ameliorate these effects.

1.5 **Objectives**
The objectives are to determine the:

- IC\textsubscript{50} following exposure to a range of FB\textsubscript{1} and allicin concentrations respectively
- involvement of oxidative stress as a mechanism of FB\textsubscript{1}-mediated cytotoxicity, and allicin response to the cytotoxic manifestations
- effect of FB\textsubscript{1}, allicin and a combination of FB\textsubscript{1} and allicin exposure on apoptosis induction
CHAPTER 2: LITERATURE REVIEW

2.1 Fumonisin B₁

2.1.1 Background
Fumonisins are mycotoxins produced as secondary by-products of Fusarium species (Seefelder et al., 2003). Fusarium verticillioides (Sacc.) Nirenberg (previously known as Fusarium moniliforme) and Fusarium proliferatum are two of the most prominent Fusarium species that produce fumonisins. Due to their geographical distribution and high production levels (Abuja and Albertini, 2001), these contaminating species make fumonisin a worldwide contaminant of maize, a dietary staple for human food and animal feed (Hendricks, 1999, Hassan et al., 2015).

Twenty eight types of fumonisins have been discovered and characterised into 4 groups namely, fumonisin A, B, C and P (FA, FB, FC and FP series) (Abuja and Albertini, 2001). The FB series is the most common of all the fumonisins and within it, FB₁ is the most notorious in prevalence and toxicity.

2.1.2 Structural composition
FB₁ (Figure 2.1) is composed of two units of propane-1, 2, 3-tricarboxylic acid (TCA) that is esterified to a 2S-amino-12S,16R-dimethyl-3S,5R,10R,14S,15R-pentahydroxy-eicosane backbone by a diester bridge at the C-14 and C-15 hydroxy groups. The presence of the four free carboxyl groups, the hydroxyl groups and the amino group make FB₁ soluble in polar solvents and insoluble in some organic solvents (Chu et al., 2012).
2.1.3 Mechanism of action

Fumonisin B₁ is similar in structure to the bioactive lipid signaling molecules, sphingolipids (Figure 2.1). Sphingolipid biosynthesis is a de novo pathway set off by the condensation of serine and palmitoyl-CoA catalysed by serine palmitoyl transferase to form 3-keto-dihydrosphingosine. 3-keto-dihydrosphingosine is then reduced to sphinganine (dihydrosphingosine). The sphinganine is subsequently N-acylated by dihydroceramide synthases to generate dihydroceramide which is desaturated creating a 4, 5-trans-double bond to form ceramide. This pathway produces and culminates in the production of bioactive intermediates comprising sphingosine, dihydroceramide and ceramide (Wang et al., 1991, Merrill, 2002).

A study by Wang et al. (1991) found that fumonisins could disrupt the de novo metabolism of sphingolipids (Figure 2.2) via the inhibition of ceramide synthase (Wang et al., 1991). The consequence of FB₁-induced inhibition of ceramide synthase in vivo is the accumulation of sphinganine, an occasional increase of sphingosine, high sphinganine/sphingosine ratio,
accumulation of 1-phosphate metabolites of sphinganine and sphingosine and low complex sphingolipids (Voss and Riley, 2013). These changes represent an important step in fumonisin toxicity that sets off a series of reactions which disrupt cell growth, differentiation and cell injury \textit{in vitro} and \textit{in vivo}.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure2.2.png}
\caption{Schematic representation of sphingolipid biosynthesis in animal cells and the points at which FB\textsubscript{1} inhibits ceramide synthase (Merrill Jr et al., 2001, Stockmann-Juvala and Savolainen, 2008).}
\end{figure}

An alternative mechanism of action was recently described in a study by Chuturgoon \textit{et al.} (2014). this study associated FB\textsubscript{1} with DNA methylation, a major epigenetic modification (Chuturgoon \textit{et al.}, 2014).
2.1.4 Epigenetics

Epigenetic modifications are heritable changes in gene activity or function that are not encoded by the DNA sequence. Major components of epigenetic modification include DNA methylation, histone modification and microRNAs (miRNAs) which contribute to gene expression through altering the chromatin structure and DNA accessibility. DNA methylation and histone modification exert their actions at the transcriptional level and miRNAs at the post-transcriptional level (Baccarelli and Bollati, 2009, Halušková, 2010, Sharma et al., 2010).

2.1.5 Effects of fumonisin B₁ in animals

The effects of FB₁ exposure have continuously proven to vary in different species (Stockmann-Juvala and Savolainen, 2008, Chuturgoon et al., 2014). Reports have shown FB₁ to induce leukoencephalomalacia (LEM) in horses (Marasas et al., 1988, Kellerman et al., 1990, Marasas et al., 2014), pulmonary oedema in swine (Harrison et al., 1990) and OC in humans (Sydenham et al., 1990, Rheeder et al., 1992, Yoshizawa et al., 1994, Sun et al., 2007) FB₁ is also renowned for its nephrotoxic, hepatotoxic and neurotoxic effects in laboratory animals (Riley et al., 1994, Voss et al., 1995, Gelderblom et al., 2001, Marasas et al., 2004).

The presence of FB₁ in maize and wheat products intended for human consumption has been linked to human health risks including OC and neural tube defects (NTD). OC has been epidemiologically linked to FB₁ contaminated maize in studies done in South Africa where reports showed that a high risk of OC development was proportional to an increased intake of maize (Marasas and WFO, 1988).

Neural tube defects are congenital malformations of the brain and spinal cord that occur when the embryonic neural tube fails to close. This common birth defect has been associated with high exposure of FB₁ in areas known to have high consumption of maize. A hypothesis for this association stated that FB₁-induced inhibition of ceramide synthase that disrupts sphingolipid biosynthesis and
subsequently folate transport causes a decrease in folate uptake that increases the risk of NTD (Hendricks, 1999, Marasas et al., 2004, Missmer et al., 2005, Stockmann-Juvala and Savolainen, 2008).

Many studies have been done on the effects of FB1 toxicity in animals, commonly laboratory animals. These effects include hepatotoxicity, nephrotoxicity, immunotoxicity, neurotoxicity and pulmonotoxicity. Studies have demonstrated the occurrence of FB1-induced hepatotoxicity and nephrotoxicity in laboratory animals through repeated exposure. Gelderblom et al. (1988) found that male BD IX rats developed chronic toxic hepatosis when treated with 48 mgFB1/kg body weight daily (12 days) then 70 mgFB1/kg body weight daily (9 days) in a 21-day experiment (Gelderblom et al., 1988). In a study by Voss et al. (1993) both hepatotoxicity and nephrotoxicity were demonstrated in a study done on Sprague-Dawley rats (Voss et al., 1993). FB1-induced toxicity in the liver and kidneys is a common consequence of various feeding studies (Voss et al., 1993, Riley et al., 1994, Howard et al., 2001).

The immunological effects of FB1 have only been studied based on the cytokine profile of different cell types and organs. A study on Sprague-Dawley rats demonstrated that FB1 affected the humoral immune response of the male rats (Tryphonas et al., 1997). The most common neurotoxic effect of FB1, LEM developed through oral or intravenous administration was found in horses (Kellerman et al., 1990).

FB1 has been shown to exhibit carcinogenic effects on various cells in animals and humans. The International Agency for Research on Cancer (IARC) has classified FB1 as a type 2B carcinogen, a possible human carcinogen. A 2-year study by Howard et al. (2001) found FB1 to be a renal carcinogen causing renal tubule tumours in male F344 rats at doses 50 and 150 ppm and a hepatocarcinogen inducing hepatocellular adenomas and carcinomas in female B6C3F1 mice at doses
50 and 80 ppm (Howard et al., 2001). In rat kidney epithelial cells, FB₁ was reported to have altered global histone modification (Sancak and Ozden, 2015) and in human hepatoma cells, found to have induced global DNA methylation (Chuturgoo et al., 2014) which may be linked to the progression of cancer, tumourigenesis or cell cycle arrest.

2.1.6 Molecular effects of FB₁ exposure
The two main consequences of FB₁ exposure are oxidative stress and apoptosis. Together, these processes are responsible for the diverse effects associated with FB₁ exposure.

2.1.6.1 Oxidative Stress
Reactive oxygen species (ROS) are by-products of aerobic respiration in the process of normal cellular metabolism (Held, 2012). Oxygen is converted through a series of reduction reactions to superoxide (O₂⁻), hydrogen peroxide (H₂O₂) and water (H₂O) (Figure 2.3). These radical oxygen species are counteracted by an arsenal of antioxidant proteins or enzymes. Superoxide dismutases (SODs) are known to be the major antioxidant defense proteins, catalysing the conversion of O₂⁻ into H₂O₂– in the mitochondria, SOD2 (MnSOD) is the isoform responsible for this reaction. The resulting H₂O₂ is further converted to H₂O, catalysed by glutathione peroxidase (GPx), catalase or peroxiredoxin (Prxs) (Figure 2.3). GPx simultaneously catalyses the formation of oxidised glutathione (GSSG) from reduced glutathione (GSH) (Figure 2.3) (Halliwell, 2007, Gupta et al., 2014). Glutathione is an important antioxidant vastly populated in the liver and is needed by the kidneys to maintain homeostasis, making it crucial for the body’s detoxification process (Lash, 2005, Zitka et al., 2012). Reduced GSH is a ROS scavenger. On DNA and other biomolecules it demolishes oxygen free radicals, radical centers and reactive hydroxyl free radicals (Mytilineou et al., 2002). The relative concentration of these antioxidants is controlled by the nuclear factor erythroid 2 (NFE2)-related factor 2 (Nrf2) (Fukai and Ushio-Fukai, 2011), a transcription factor that exerts cytoprotective properties against oxidative stress.
Intracellular and/or extracellular ROS are essential for biochemical processes such as cellular function, intracellular signaling and defense against microorganisms at low levels (Fukai and Ushio-Fukai, 2011). However, at increased levels ROS inactivate or deplete antioxidant enzymes (Lushchak, 2015), resulting in an accumulation or inadequate removal of ROS and oxidative stress (Fukai and Ushio-Fukai, 2011). Oxidative stress therefore is increased toxic levels of ROS production surpassing the cell’s ability to counteract the effects. The induction of oxidative stress not only damages tissues, but causes genetic instability and disrupts normal cellular functioning (Towner et al., 2003, Mehta et al., 2007, Matur et al., 2011, Abbes et al., 2016, Khan et al., 2018b).

**Figure 2.3**: Schematic representation of the synthesis and detoxification of ROS (Merksamer et al., 2013).

FB$_1$-induced oxidative stress varies from cell type to host species, demonstrated by studies done on OC cells, mice, hens, humans and rats (Towner et al., 2003, Mehta et al., 2007, Matur et al., 2011, Abbes et al., 2016, Khan et al., 2018b). In humans, it is postulated that oxidative stress mediates the
immunotoxic effects of FB$_1$ (Stockmann-Juvala et al., 2007, Klaric et al., 2008, Domijan et al., 2012). The findings of a study by Abbes et al. (2016) further verified the role of FB$_1$ in inducing oxidative stress by demonstrating increased levels of oxidative stress markers in mice spleen where ROS production was increased overall. It hypothesised that genotoxicity or mutagenicity may be indirect consequences of FB$_1$-induced oxidative damage (Hassan et al., 2015, Abbes et al., 2016). It must also be noted that cells may respond to oxidative stress by triggering apoptosis.

2.1.6.2 Apoptosis

The regulation of homeostasis and other cellular processes helps maintain cellular functionality. Apoptosis is a biological process of programmed cell death marked by cell shrinkage, DNA fragmentation and chromatin condensation that occurs as a mechanism of defense that removes damaged cells (Kerr et al., 1972, Renehan et al., 2001). This process is controlled by caspases, which are initially produced as inactive pre-enzymes (procaspases) before activation through different cleavage processes at specific aspartate residues (Shi, 2004, Elmore, 2007).

Apoptosis occurs via the intrinsic and extrinsic pathways (Figure 2.4). The intrinsic pathway is triggered by a number of mitochondrial stimuli such as DNA damage, oxidative stress and growth factor deprivation. These stimuli produce signals intracellularly mediated by the mitochondria. The extrinsic pathway is activated through the binding of death receptors to ligands to transmit apoptotic signals (Elmore, 2007, Mukhopadhyay et al., 2014).

The intrinsic pathway, also known as the mitochondrial pathway is mainly regulated by the Bcl-2 family proteins responsible for mitochondrial membrane permeability (Figure 2.4). They stimulate BH3 family proteins that activate BAX and BAK pro-apoptotic effectors. The process is triggered by several stimulating factors that disturb the electrochemical gradient, increase membrane permeability
and release apoptogenic factors. A disturbance in the mitochondrial membrane integrity results in either necrosis and/or the release of cytochrome c into the cytosol which activates initiator caspase 9 through the assembly of an apoptosome (a multiprotein caspase-activating complex comprised of Apaf1 and procaspase 9) that activates effector caspases 3/7 that function to execute apoptosis (Hassan et al., 2014, Baig et al., 2017).

The death receptor-mediated pathway is triggered when the signaling mechanism of death domain (DD)-containing tumour necrosis factor (TNF) family receptors activates caspases (Figure 2.4). Death receptor ligands bind to the plasma membrane forming a death-inducing signaling complex (DISC) to activate procaspases 8 and 10. Caspase 8 which contains death effector domains on the N-terminal prodomains initiates proapoptotic cascade of caspases. Caspase 8 can either activate the intrinsic pathway through the activation of tBID which triggers the oligomerisation of Bak to release cytochrome c or activates caspase 3 (Figure 2.4). Caspase 3 cleaves inhibitor of caspase activated DNase (iCAD) to CAD that leads to DNA fragmentation, apoptotic blebs, apoptotic bodies and chromatin condensation (Hassan et al., 2014, Koff et al., 2015).
Exposure to FB₁ results in organ-specific toxic manifestations. Likewise, the induction of apoptosis by FB₁ is dependent on cell type and species, as well as the concentration (Khan et al., 2018). For example, neuroblastoma cells are sensitive to FB₁-induced apoptosis (Stockman-Juvala, 2006), while apoptosis is inhibited in HepG2 cells (Chuturgoon et al., 2015). A number of animal studies investigating the consequences of FB₁ exposure describe one of its effects to be apoptosis. This effect results from FB₁-induced disruption of sphingolipid metabolism by inhibiting the enzyme ceramide synthase (Wang et al., 1991, Brenner and Kroemer, 2000, Minervini et al., 2004). Dragan et al. (2001) characterised FB₁-induced apoptosis as mechanism of action in carcinogenesis (Dragan et al., 2001).
2.2 Medicinal plants

Plants have been used for medicinal purposes for many years. Scientific research as shown their health benefits to span from antiseptic properties to the treatment of cancer (Greenwell and Rahman, 2015). Their wide array of health benefits has since prompted the identification of plant species and their extracts for the development of some therapeutic drugs. Among many, *Allium sativum* and its constituents is one of the most commonly studied plants for its potential to alleviate and prevent disease (Amagase, 2006, Iciek *et al.*, 2009).

2.2.1 *Allium sativum*

*Allium sativum*, more commonly known as garlic, is a medicinal plant whose use in different cultures has had many beneficial health effects through history. Its biological benefits include anti-cancer properties, enhanced immune function, reduced cardiovascular disease risk factors, anti-bacterial, anti-fungal and anti-parasitic properties among many (Mikaili *et al.*, 2013, Gruhlke *et al.*, 2016). The varying effects of garlic have been attributed to its different preparations under four categories namely garlic oil, aged garlic extract (AGE), garlic powder and garlic oil macerate. A common characteristic of garlic is its distinct smell caused by oil-soluble sulfur constituents; whole intact garlic comprises primary sulfur-containing molecules which include $\gamma$-glutamyl-$S$-alk(en)yl-L-cysteines and $S$-alk(en)yl-L-cysteine sulfoxides (i.e alliin) (Bhandari, 2012)

2.2.2 Allicin, an organosulphur compound from garlic

When the plant tissue is damaged or whole garlic is macerated, garlic compounds undergo rapid conversion into organosulphur compounds such as allicin (alkyl alkane-thiosulfinate) which represents the biologically active component of crushed garlic. Allicin is synthesised via a two-step reaction that begins when the enzyme alliinase that hydrolyses alliin within the cytosol to produce dehydroalanine and odiferous sulfenic acid (Bat-Chen *et al.*, 2010). A spontaneous condensation reaction of two allyl sulfenic molecules (2-propenesulfenic acid) occur forming one allicin molecule.
Allicin may then be degraded to ajoene by heating, or condense with other allicin molecules yielding secondary products including diallyl sulfide (DAS), diallyl disulfide (DADS) and diallyl trisulfide (DAT) and dithiins (Figure 2.5). In a different pathway, γ-glutamyl-cysteines are converted to S-allylcysteine (SAC) (Mikaili et al., 2013, Luo et al., 2016).

**Figure 2.5:** The two-step biosynthesis of allicin involves the enzymatic conversion of alliin to 2-propenosulfenic acid, and subsequent condensation to allicin. Secondary products are formed by various means. Adapted from (Gruhlke et al., 2016, Luo et al., 2016).

### 2.2.3 Effects of allicin

Allicin has been vastly studied for its many health benefits including anti-cancer activity through allicin-induced apoptosis and inhibition of cancer cell proliferation in different cell lines (Siegers et al., 1999, Hirsch et al., 2000, Bat-Chen et al., 2010). Allicin-induced apoptosis varied in different cell lines, and was found to be both caspase-dependent (Oommen et al., 2004, Bat-Chen et al., 2010) and caspase-independent (Park et al., 2005). It has been suggested that inhibition of cell proliferation
mediated by allicin may be associated with modulation of tubulin, which interferes with mitotic spindle formation and cell division (Prager-Khoutorsky et al., 2007).

The exact mechanisms by which allicin induces its antioxidant effects remains unclear. However, studies have shown it to have free radical scavenging effects, ability to inhibit the production of $O_2^-$ and react with thiol-containing compounds. This reactive sulfur species also has the ability to oxidise GSH and cysteine residues in various proteins because of its membrane permeability. Studies have also shown that in the presence of ROS, allicin induces phase II detoxification and upregulates nuclear factor erythroid-related factor 2 (Nrf2), thereby fulfilling an essential role in the replenishment of several antioxidants, including SOD2 and GSH (Gruhlke et al., 2016).
3.1 Materials

FB₁ was purchased from Sigma Aldrich (Johannesburg, South Africa (SA)). The human embryonic kidney (Hek293) cells were purchased from Highveld Biological (Johannesburg, SA). Cell culture reagents were acquired from Whitehead Scientific (Johannesburg, SA). Western blotting reagents were purchased from Bio-Rad (Hercules, CA). Antibodies and Promega kits were bought from Anatech (Johannesburg, SA). Other reagents were obtained from Merck (Darmstadt, Germany).

3.2 FB₁ and allicin preparation

Stock solutions of FB₁ (1468.5 µM) and allicin (1848.8 µM) were prepared in 0.1 M phosphate buffered saline (PBS, pH 7.4). These stocks were stored at 4 °C and diluted for subsequent assays.

3.3 Cell culture

The Hek293 cell line was reconstituted by transferring a vial of cryopreserved cells into a sterile 25 cm³ cell culture flask containing 10 ml of complete culture medium (CCM) comprising of Dulbecco’s Modified Eagle’s Medium (DMEM) supplemented with 10 % foetal calf serum, 25 mM HEPES, 1 % L-glutamine and 1 % penicillin-streptomycin-fungizone. The flask was incubated for 4 hours (4 h) at 37 °C, in a 5 % CO₂-supplemented incubator, after which the CCM was discarded and replaced with 5 ml of fresh CCM. Cells were maintained until 90 % confluence was reached. Confluent flasks were washed with phosphate-buffered saline (PBS); the cells were dislodged and counted (150 µl of CCM, 50 µl of trypan blue, 50 µl of cell suspension in an eppendorf and transferring 10 µl of this mixture onto a haemocytometer) for sub-culturing or storage.
3.4 Methylthiazol tetrazolium (MTT) Assay

3.4.1 Principle

The methylthiazol tetrazolium (MTT) assay was used to measure cell viability and cytotoxicity in Hek293 cells following exposure to FB₁ and allicin respectively. The yellow MTT salt that crosses the membranes of living cells is reduced to a purple formazan product by mitochondrial succinate dehydrogenase (SDH) (Figure 3.1) and therefore reflects the number of viable cells (Abate et al., 1998). The MTT assay generates a half maximum inhibitory concentration (IC₅₀) that refers to a concentration that inhibits a population’s biochemical functioning by 50 %. The IC₅₀ will be used as the treatment dose in subsequent assays.

![Figure 3.1: Metabolic cellular reduction of yellow MTT salt into a purple formazan (Riss et al., 2004).](image-url)
3.4.2 Protocol

The Hek293 cells were seeded into a 96-well microtitre plate at a density of 15,000 cells/well (200 µl) in triplicate and allowed to attach overnight. Cells were incubated with a range of FB1 and allicin concentrations (0-1000 µM) at 37 °C for 24 h. Thereafter, the cells were incubated with 20 µl MTT salt solution (5 mg/ml in 0.1 M PBS) and 100 µl CCM at 37 °C. After 4 h incubation, the MTT salt was discarded and replaced with 100 µl of DMSO for the solubilisation of formazan crystals and further incubated for an hour at 37 °C. The optical density of the samples was read at 570 nm and 690 nm using the Bio-Tek µQuant spectrophotometer (USA). The cell viability and the concentration dependent response curve were plotted using GraphPad Prism v5.0 software and an IC50 was generated.

Flasks of Hek293 cells were grown to confluence and treated as follows:

- control - CCM only
- IC50 of FB1
- IC50 of allicin and
- Combined IC50 of FB1 + IC50 of allicin

After 24 h treatment, the cells were washed, resuspended in PBS, counted and the cell number was adjusted for use in subsequent assays.

3.5 Bioluminescence Assays

Bioluminescence is the generation and emission of light by an organism due to a chemical reaction whereby chemical energy is transformed into light energy and the outcome is chemiluminescence (Hastings, 1968). The assay is based on the luciferin-luciferase reaction that occurs in the firefly. Luciferase is the enzyme that catalyses the monooxygenation of D-luciferin in the presence of magnesium, ATP and oxygen, producing inactive oxyluciferin. The intensity of light that is produced
by this reaction is directly proportional to the concentration of the substrate being measured (Lundin, 2000).

3.5.1  *CytoTox-ONE™* Homogenous Membrane Integrity Assay

3.5.1.1  Principle

While the MTT assay was used to estimate the number of viable cells, studies have shown that it is invaluable to measure the leakage of cytoplasmic components into the surrounding culture medium in order to quantify non-viable cells. This assay quantifies the fluorescent signal generated by resofurin when lactate dehydrogenase (LDH) released from cells with damaged membranes into the culture medium participates in a coupled reaction that results in the conversion of resazurin to resofurin (Figure 3.2) (Ronken, 2009).

![Diagram of the CytoTox-ONE assay](image)

**Figure 3.2**: Leakage of LDH from a cell with a damaged membrane (Ronken, 2009).

3.5.1.2  Protocol

The protocol was followed according to the manufacturer’s instructions. Briefly, treated Hek293 cells (2 x 10⁴ cells in 100 µl of treatment medium) were transferred to a 96-well white microtitre plate in triplicate and equilibrated to room temperature (RT). Lysis solution was added to all the wells (2 µl).
Thereafter, 100 µl of Promega CytoTox-ONE™ Reagent (#G7890) was aliquoted into all the wells and mixed well before incubation of 10 min at RT. Subsequently, 100 µl of stop solution was added into each well and the microtitre plate was put on the shaker for 10 sec. The fluorescence was obtained using the Promega GloMax®-Multi+ Detection Plate Reader (Turner Bio-systems, Sunnyvale, USA) with an excitation wavelength of 560 nm and an emission wavelength of 590 nm.

\[
\% \text{ cytotoxicity} = 100 \times \frac{(\text{experimental} - \text{culture medium background})}{(\text{max LDH release} - \text{culture medium background})}
\]

3.5.2 Caspases

3.5.2.1 Principle

The main workers of apoptosis are a family of proteins called caspases. These are cysteine-aspartate proteases that are involved in the initiation and execution of apoptosis (Garrido and Kroemer, 2004). The caspase activity is based on cleavage of the caspase-luciferin substrate by luciferase. This reaction produces light which is directly proportional to caspase activity (Figure 3.3).
3.5.2.2 Protocol

The activities of caspase 3/7, 8 and 9 were detected with the Promega Caspase-Glo® assay (#G8091, # G8201 and #G8211 respectively). Briefly, 50 μl of cell suspension (20,000 cells/well in 50 μl 0.1 M PBS) were seeded into duplicate wells of a white, opaque 96-well microtitre plate. As per manufacturer’s guidelines, caspase-Glo®-3/7 reagent was reconstituted and 50 μl was added into each well. Thereafter, the plate was incubated in the dark (30 min, RT). The procedure was completed in a similar fashion for caspase 8 and caspase 9. Luminescence was detected using a Modulus™ microplate luminometer (Turner Bio-systems, Sunnyvale, USA) as relative light units (RLU).
3.5.3 **CellTiter-Glo® Luminescent Cell Viability Assay**

3.5.3.1 **Principle**

ATP is the universal energy transducer in mammalian cells produced during cellular respiration. Levels of intracellular ATP are a useful indicator of respiratory capacity and mitochondrial function. The assay uses bioluminescence to determine intracellular ATP levels to indicate the number of viable cells in culture. During this reaction, cellular ATP is used in the luciferase reaction to generate a light signal (Figure 3.4). The intensity of light that is produced by this reaction is directly proportional to the concentration of intracellular ATP (Lundin, 2000).

![Figure 3.4](image)

**Figure 3.4**: ATP-dependant oxidation of D-luciferin to produce light (Govender, 2016).

3.5.3.2 **Protocol**

The ATP inside the cells was quantified using the Promega CellTitre-Glo® Assay (#G7570), whereby 50 µl of cell suspension (20,000 cells/well in 50 µl 0.1M PBS) was plated into a 96-well luminometer plate in duplicate. Thereafter, 50 µl of reconstituted CellTiter-Glo® substrate was added in each well. The plate was incubated in the dark for 30 min at RT. The amount of light produced was quantified using the Modulus™ microplate luminometer (Turner Bio-systems, Sunnyvale, USA) and expressed in RLU.

3.5.4 **Mitochondrial Toxicity Assay**

3.5.4.1 **Principle**

The Promega Mitochondrial ToxGlo™ Assay (#G8000) serves to predict the integrity of the mitochondria and cellular ATP levels by looking at specific biomarkers that correspond with the normal functioning and state of the mitochondria. This assay achieves such by measuring protease...
activity that shows cell membrane integrity and measures ATP levels through a luminescent signal (Figure 3.5).

![Figure 3.5: Luminometric measurement of protease activity and ATP in the mitochondria (PT. Indolab utama protocols).](image)

3.5.4.2 Protocol

To a white 96-well microtitre plate, 50 µl of Hek293 cell suspension (20,000 cells) was added into each well and the cells were allowed to attach overnight at 37 °C (5 % CO₂). Cells were treated in duplicates for 24 h; then 20 µl of 5X cytotoxicity reagent was added into each well, mixed then incubated for 30 min at 37 °C. Fluorescence was then measured at 485 nmEx/520–530 nmEm. The assay plate was then equilibrated to room temperature for 10 min before the addition of 100 µl ATP Detection Reagent to each well and mixed for 5 min. The luminescence was then measured after 1 hr.

3.5.5 Ros-Glo H₂O₂ Assay

3.5.5.1 Principle

ROS are generated from the normal metabolic processes that occur in our body. The overproduction of ROS leads to oxidative stress that causes DNA and protein damage as well as lipid peroxidation.
The $O_2^-$ is the most common source of ROS. Superoxide is converted to $H_2O_2$ by SOD. This assay serves to measure the amount of $H_2O_2$ by coupling it in a luciferase reaction where the light produced is quantified (Figure 3.6). It also directly links the amount of $H_2O_2$ to $O_2^-$ production.

**Figure 3.6**: The chemistry of ROS-Glo™ $H_2O_2$ Assay (Promega protocols).

3.5.5.2 Protocol

Hek293 cells were seeded into a 96-well plate with their respective treatments (80 µl) and allowed to attach in 24 h incubation (37 °C at 5 % CO₂). The $H_2O_2$ substrate was thawed and diluted accordingly. Thereafter, 20 µl of the $H_2O_2$ substrate was added into each well and incubated for 6 h (37 °C at 5 % CO₂). Subsequently, 100 µl of the ROS-Glo™ detection solution was pipetted into each well and incubated for 20 min at RT. The luminescence was recorded using a Modulus™ microplate luminometer (Turner Bio-systems, Sunnyvale, USA) and was analysed as RLU.
3.5.6 Glutathione Assay

3.5.6.1 Principle

Glutathione is an antioxidant that can prevent damage to vital cellular components caused by ROS such as free radicals and lipid peroxides. This assay serves to quantify GSH levels based on the conversion of a luciferin derivative into luciferin in the presence of glutathione. This conversion is catalysed by glutathione S-transferase (GST). The signal generated is proportional to the amount of glutathione present in the sample (Figure 3.7).

![Glutathione Assay Diagram](image)

Figure 3.7: GSH-Glo™ Glutathione assay overview (Promega protocols).

3.5.6.2 Protocol

In a 96-well white plate, 20,000 treated Hek293 cells were added and incubated (37 °C at 5 % CO₂). GSH standards were prepared according to the manufacturers guidelines and added to the wells in duplicate. Thereafter 100 µl of the 2X GSH-Glo™ reagent was pipetted into each well. The plate was then incubated for 30 min at RT. Thereafter, 100 µl of the Luciferin Detection reagent was added into each well and briefly mixed before an incubation of 15 min at RT. The luminescence was then measured with the use of a Modulus™ microplate luminometer (Turner Bio-systems, Sunnyvale, USA) and recorded as RLU.
3.6 Single Cell Gel Electrophoresis (SGCE) Assay

3.6.1 Principle

DNA fragmentation in individual cells is detected by the comet assay (Figure 3.8). DNA fragmentation is a late characteristic of apoptosis, a result of the chain reaction that occurs due to the caspase cascade. Healthy, undamaged DNA maintains a highly organised, supercoiled structure within the nucleus whilst damaged DNA loses its structural organisation and migrates out of the nucleus when subjected to an electric field (Godard et al., 1999). When an electric field is applied to cells embedded in agarose gel, DNA migration out of the nucleus forms a ‘comet’, where the head contains intact, undamaged DNA and the tail contains damaged, fragmented DNA. The comet assay utilises fluorescence microscopy to visualise the presence and extent of DNA damage (Singh et al., 1988).

![Diagram](image)

**Figure 3.8**: An overview of the comet assay process (Sigma-aldrich protocol).
3.6.2 Protocol

Frosted end microscope slides were used for the SCGE assay (2 for each treatment). The slides were prepared with three layers: the first layer consisted of 2 % low melting point agarose (LMPA, 800 µl) covered with a coverslip at the top of the gel and was set at 4 °C for 10 min; the second layer consisted of 1 % LMPA (400 µl) with Gel Red (1 µl) and cell suspension (20,000 cells in 30 µl 0.1 M PBS) covered with a coverslip and solidified at 4 °C for 10 min; and the third and last layer consisted of 1 % LMPA (400 µl) covered with a coverslip at the top, which was left to set at 4 °C for 10 min. The same procedure was repeated for all the slides.

Coverslips were removed and solidified gels were submerged in cold cell lysis buffer [100 mM EDTA, 2.5 M NaCl, 1 % Triton X-100, 10 % DMSO, and 10 mM Tris (pH 10)] for 1 h at 4 °C. Thereafter, the slides were submerged in electrophoresis buffer [1 mM Na₂ EDTA (pH 13) and 300 mM NaOH] for 20 min at RT to allow for equilibration prior to electrophoresis (25 V, 35 min at room temperature) using a Bio-Rad compact power supply.

The slides were rinsed three times (5 min each) in (0.4 M Tris, pH 7.4) to neutralise the samples prior to replacement of coverslips. The slides were then viewed using a fluorescent microscope (Olympus IX51 inverted microscope, excitation: 510-560 nm; emission 590 nm). Images of 50 cells and comets were captured from each slide. The comet tail lengths were quantified using a Soft Imaging system and reported as tail lengths in µm.

3.7 Thiobarbituric acid reactive substances (TBARS) Assay

3.7.1 Principle

The detection of oxidative stress has relied largely on the quantification of compounds such as malondialdehyde (MDA), which are formed by degradation of initial products of free radical attack on membrane lipids (Oakes and Van Der Kraak, 2003). The reaction of MDA with 2-thiobarbituric
acid (TBA) produces a coloured product that can be detected at 532 nm (Figure 3.9). This assay is therefore one of the most widely used estimators of oxidative stress that measures MDA production, an indication of lipid peroxidation induced by oxidative stress.

![Diagram of MDA/TBA compound formation](image)

**Figure 3.9**: Formation of MDA /TBA compound via the TBARS assay [Adapted from R&D systems protocol].

### 3.7.2 Protocol

The supernatants of each treatment (200 μl) were added into appropriately labelled, clean glass test tubes in duplicates containing 200 μl of 2% phosphoric acid (H₃PO₄). A positive (containing MDA) and negative control (CCM only) was included in this experiment. Thereafter, 400 μl of TBA/Butylated hydroxytoluene (BHT) solution was added to every sample except the negative control, this tube received 400 μl of 3 mM HCl.

Each test tube was briefly vortexed, and the pH of each sample was adjusted to 1.5 using 1 M HCl. The tubes were subsequently boiled in a water bath for 15 min at 100 °C to allow for optimal hydrolysis of MDA-adducts. After the samples cooled to RT, 1500 μl of butanol was added to each tube to extract MDA by separating the solution into two phases. The tubes were then vortexed and allowed to stand for the distinction of the two phases.
Lastly 200 μl of the upper butanol phase in each test tube was transferred to a 96-well microtitre plate in duplicates. The absorbance was measured at 532 nm with a reference wavelength of 600 nm using a Bio-Tek μQuant spectrophotometer (USA).

3.8 Nitric Oxide Assay

3.8.1 Principle

Reactive nitrogen species (RNS) are formed when nitric oxide (NO) reacts with superoxide to produce peroxynitrite. It is therefore important to evaluate the production of NO to determine if nitrosative stress is present. The assay indirectly quantifies NO and therefore gives a measure of the RNS present.

The nitric oxide assay is a traditional method used to measure the conversion of L-[3H] arginine to L-[3H]citrulline. However, because nitric oxide is oxidised to nitrates and nitrites in biological samples, it is these end-products that are quantified in this two-step reaction assay used to measure oxidative stress. The Greiss reaction is used as follows: vanadium (III) chloride (VCl₃), sulphanilamide (SULF) and N-1-Naphthyl ethylenediamine dihydrochloride (NEDD) are each added in quick succession to facilitate reduction, inhibition of enzymatic reactions involving para-aminobenzoic acid and colour change (detected at 540 nm) respectively (Figure 3.10) (Beda and Nedospasov, 2005).
Figure 3.10: Nitric oxide assay represented by a two-step azotisation reaction that produces a diazo product (Bryan and Grisham, 2007).

3.8.2 Protocol

Sodium nitrate standards were prepared with a concentration range of 0–200 µM and 50 µl of each standard as well as the respective treatments were plated into a 96 well plate in duplicate. Subsequently, 50 µl VCl₃ and 25 µl SULF were added in all the wells (standards and samples) followed by the addition of 50 µl NEDD. The plate was then incubated at 37 °C for 45 min. Following incubation, the plate was read and quantified using a Bio-Tek µQuant plate reader (USA) at a wavelength of 540 nm and a reference wavelength of 690 nm. A standard curve using the standards was then plotted and the concentration of RNS of the samples was extrapolated from the graph.
3.9 Western Blotting

3.9.1 Principle

Western blotting is an analytical technique used to detect the proteins in a homogenous sample. It relies on the principle that the proteins are separated through a sodium dodecyl sulfate (SDS)-Polyacrylamide gel using an electric field (Figure 3.11). The extent of migration is determined by the size of the proteins. The proteins are then transferred onto a nitrocellulose membrane and immunoprobing takes place for the actual western blot (Figure 3.12). Proteins are represented as bands at the site of the antigen-antibody reaction and the band width is proportional to the quantity of the respective protein (Mahmood and Yang, 2012). Antioxidant and apoptotic proteins were quantified via western blotting.

3.9.2 Protocol

Crude protein was isolated from treated cells using Cytobuster™ reagent containing protease and phosphatase inhibitors (300 µl). Flasks were placed on ice for 15 min, and then cells were scraped off the flasks and transferred to 1.5 ml eppendorf tubes which were also placed on ice for a further 15 min. The cells were then centrifuged at 2 000 x g for 5 min (4 °C) to remove cellular debris. The crude protein obtained was quantified using the bichinchoninic acid (BCA) assay.

Bovine serum albumin (BSA) standards (0–1 mg/ml) were prepared to determine the concentration of the protein samples. The standards and samples were then pipetted (25 µl) in duplicate into a 96-well microtiter plate. The BCA reagent (198 µl BCA: 4 µl CuSO₄) was added in each well (200 µl) and incubated at 37 °C for 30 min. The absorbance was detected using Bio-Tek µQuant spectrophotometer (USA) at 562 nm. A standard curve was constructed and proteins were standardised to 1 mg/ml. Laemmli buffer (dH₂O, 0.5 M Tris-HCl (pH 6.8), glycerol, 10 % SDS, β-mercaptoethanol, 1 % bromophenol blue) was added to each sample (1:4, total volume 250 µl), and heated to 100 °C (5 min).
**Figure 3.11**: Western blot procedure from sample preparation to protein electrophoresis [Adapted from Creative Diagnostics protocol].

The standardised, denatured proteins (25 µl) were loaded onto a SDS-polyacrylamide gel (10 % resolving gel and 4 % stacking gel) to separate proteins (1.5 h at 150 V) using running buffer (Tris, glycine, SDS, dH2O), after which transfer onto nitrocellulose membranes was achieved with transfer buffer (Tris, glycine, methanol, dH2O) using the Transblot®Turbo™ Transfer system (Bio-Rad). Membranes were blocked with 5 % BSA in Tween20 Tris-buffered saline (TTBS, 25 mM Tris (pH 7.5) 150 mM NaCl, 0.05 % Tween 20) for 2 h, and incubated with primary antibodies (GPx1 [#3286], SOD2 [#13141], Nrf2 [#12721], p53 [#48818], Bax [#5023], cPARP [#9542], catalase [#12980] and Hsp70 [#46477]) in 5 % BSA/TTBS (1:1000 dilution) overnight. The primary antibody was removed and membranes were washed five times with TTBS (10 min each), then incubated in secondary antibody (anti-rabbit IgG [#7074] or anti-mouse IgG [#70746]) as required, in 5 % BSA/TTBS (1:2500 dilution) for 2 h. Following incubation, membranes were washed with TTBS (5x 10 min each). Clarity Western ECL Substrate (Bio-Rad) (150 µl) was added to the membranes and images were captured using a Molecular Imager® Chemidoc™ XRS and BioRad imaging system and actin [#12141] as a loading control.
Figure 3.12: Representation of immunoblotting illustrating the binding of a primary antibody onto the target protein, followed by the secondary antibody-enzyme conjugate. This then allows for the emission of a chemiluminescent detection signal (elabscience protocols).

3.10 Quantitative- Polymerase Chain Reactions (qPCR)

3.10.1 Principle

Quantitative- Polymerase Chain Reactions (qPCR) also known as real time PCR, is a reliable technique that both detects and measures products of the PCR process. When the PCR product is directly proportional to the amount of template DNA from the beginning of the PCR process, data is then collected in the exponential growth stage (Arya et al., 2005). The qPCR process occurs in two steps, first mRNA is isolated from cells, and then the mRNA is reverse transcribed to single-stranded complementary DNA (cDNA). Furthermore, qPCR is characterised by the addition of a fluorescent dye, SYBR green to the master mix (SYBR green, nuclease-free water, forward and reverse primer) (Hernandez-Rodriguez and Ramirez, 2012).
3.10.2 Protocol

Cells with the relevant treatments were washed once with 0.1 M PBS. The total RNA was isolated in a series of steps. Briefly, 500 μl PBS and 500 μl Triazol was added to the flask and incubated at RT for 5 min. Thereafter, the flasks were scraped and the contents were transferred to 1.5 ml micro-centrifuge tubes and stored at -80 °C overnight. Thereafter, 100 μl chloroform was added to the thawed samples and mixed vigorously for 15 sec. The samples were incubated at room temperature for 3 min. The micro-centrifuge tubes were then centrifuged (12000 x g, 4 °C for 15 min), and the upper aqueous layer was transferred to a new 1.5 ml micro-centrifuge tube, which was placed on ice. Subsequently, 250 μl isopropanol was added to each sample and incubated on ice for 2 h before being returned to -80 °C overnight. Samples were thawed again and centrifuged (12000 x g, 4 °C for 20 min). The resulting pellet contained the total RNA and was washed with 500 μl cold 75 % ethanol. Samples were then centrifuged again (7400 x g, 4 °C for 15 min). All ethanol was discarded and the pellet was left to air-dry, before re-suspending in 15 μl RNase-free water. RNA samples were incubated at RT (3 min). A Thermo Scientific™ Nanodrop 2000 (Thermo Fisher Scientific; Johannesburg, South Africa) machine was used to determine the quality and quantity of the isolated RNA. The absorbance ratio of 260 nm/280 nm was used to determine purity of samples (~2). RNA samples were standardised to a concentration of 1000 ng/μl with RNase-free water and stored at -80 °C until required.

RNA was reverse transcribed to cDNA as per manufacturers’ guidelines using the iScript™ cDNA Synthesis kit (Bio-Rad; 107-8890): a 20 μl reaction volume containing 1 μl iScript™ reverse transcriptase, 4 μl 5X iScript™ reaction mix and the RNA template (2000 ng) made up in nuclease free water. Thermocycler conditions were set to 25 °C for 5 min, 42 °C for 30 min, 85 °C for 5 min and a final hold at 4 °C. The cDNA was diluted 1:4 to obtain a final volume of 100 μl.
Gene expression was analysed using the iQ™ SYBR Green Supermix (Bio-Rad; 170-880), according to the manufacturer’s instructions. Briefly, 1.5 μl cDNA template, 1 μl sense primer, 1 μl antisense primer, 5x iScript reaction mix and nuclease free water was made up to a reaction volume of 25 μl. The gene of interest was SOD2 [forward 5’-CAGCCCAGCCTGCTAGACGG-3’; reverse 5’-GCGTTGATGTGAGGTTCCAG-3’ (57 °C)]. All primers were obtained from Inqaba Biotechnologies (Pretoria, South Africa).

The qPCR experiments were conducted using the CFX Touch™ Real Time PCR Detection System (Bio-Rad; Hercules, California, United States). The assay was run with three replicates per treatment. GAPDH [forward 5’-CGTGAAGGACTCATGACCA-3’; reverse 5’-GCCCACGACAGTTTC-3’ (57 °C)] was used as a housekeeping gene and was amplified simultaneously under the same conditions as the treatment samples. The reaction was subjected to an initial denaturation (95 °C, 4 min), followed by 37 denaturation cycles (95 °C, 15 sec), an annealing phase (57 °C, 40 sec), and an extension phase (72 °C, 30 sec).

The acquired data was analysed using the method described by Livak and Schmittgen (2001) represented as fold-change in mRNA expression ($2^{\Delta\Delta CT}$) relative to the control (Livak and Schmittgen, 2001).

3.11 Statistical analysis

The data acquired was expressed as mean optical density with standard deviations for the MTT assay and IC₅₀ was determined using the dose-response inhibition equation (Log inhibition versus variable slope). For the data obtained from subsequent assays, the One-way ANOVA and an unpaired t-test with Welch’s correction was performed. The data was represented as means ± standard deviations. A p value of 0.05 was considered to be statistically significant. These statistical tests are available on GraphPad prism version 5.0 software.
CHAPTER 4: RESULTS

The results obtained from the experiments elaborated in Chapter 3 will be outlined. Each reported result is accompanied by a graphical illustration that shows the major changes that occurred following treatment with FB₁, allicin and a combination of FB₁ and allicin.

4.1 Methyl-thiazol-tetrazolium (MTT) Assay

The dose-response of FB₁ and allicin was assessed using the MTT assay and generated the IC₅₀ for use in this study. The cell viability decreased to 76 % at 50 µM FB₁, 72 % at 125 µM FB₁ and 29 % at 250 µM FB₁; yielding an IC₅₀ of 215 µM FB₁ (Figure 4.1A). Allicin induced a decrease in cell viability to consistently less than 30 % from 5 – 150 µM, and an IC₅₀ of 3.905 µM was derived (Figure 4.1B).

![Figure 4.1](image)

**Figure 4.1**: The percent cell viability of Hek293 cells treated with varying concentrations of FB₁ (A) and allicin (B) respectively was decreased with increasing doses when compared to an untreated control. The curves were used to calculate the IC₅₀ of both exposures.

4.2 CytoTox-ONE™ Homogeneous Membrane Integrity Assay

The CytoTox-ONE™ homogeneous membrane integrity assay was used to measure the release of LDH from cells as an indicator for cytotoxicity (Figure 4.2). Basal LDH level in untreated cells was 16700 ± 0.002983 RLU. Individual FB₁ and allicin treated Hek293 cells showed a slight reduction in
cell membrane damage, whilst LDH release in combined FB₁+allicin treated cells increased to 35150 ± 6450 RLU, indicating significant cell membrane damage and cytotoxicity.

![Bar graph showing luminescence (RLU) for different treatments of HEK293 cells.](Figure 4.2)

**Figure 4.2:** Slightly decreased LDH levels in individual FB₁ and allicin treated cells. LDH was significantly increased to 35150 ± 6450 RLU in combined FB₁+allicin treated cells. One-way ANOVA with Tukeys post-test – p = 0.0015.

### 4.3 Mitochondrial ToxGlo™ Assay

The mitochondrial integrity was assessed by measuring protease activity (Figure 4.3). There was a decrease in the mitochondrial integrity of FB₁ treated cells, while a 1.65-fold increase occurred in allicin treated cells and combined FB₁+allicin treatment restored mitochondrial integrity to control levels (313.0 ± 100.6).
Figure 4.3: Decrease in mitochondrial integrity of FB1 treated cells ($p = 0.7102$), an increase in allicin treated cells ($p = 0.3797$) and a slight decrease in FB1+allicin treated cells compared to the control.

4.4 Adenosine Triphosphate (ATP) Assay

The ATP concentration in Hek293 cells was assessed using luminometry. There was a significant increase in ATP levels for all treatments (Figure 4.4). FB1 exposure resulted in a 50 % increase in ATP concentration ($p = 0.0003$). Allicin caused a slightly greater increase of 1.62-fold ($p = 0.0024$), while a significant 1.69-fold increase was observed in the combined FB1+allicin treatment ($p < 0.0001$) when compared to the control (8989000 ± 76750).
Figure 4.4: Significantly increased ATP levels in all HEK293 cell treatments relative to the control (8989000 ± 76750).

4.5 Free radicals

4.5.1 Reactive oxygen species

This assay measures the amount of H$_2$O$_2$ in the cell. The results demonstrate increased levels of H$_2$O$_2$ in all treatments (Figure 4.5A) with a 1.43-fold ($p = 0.0671$) in FB$_1$, 1.06-fold in allicin ($p = 0.5369$) and 1.66-fold in FB$_1$+allicin ($p = 0.0002$) treated cells compared to the control (63430 ± 4343).
**Figure 4.5A**: Increased H$_2$O$_2$ levels in all HEK293 cell treatments relative to the control (63430 ± 4343).

### 4.5.2 Nitric Oxide Assay (NOS)

This assay measures the production of RNS. All three treatments showed a slight decrease in RNS formation (Figure 4.5B). Treatment with FB$_1$ causing a 33% decrease in RNS ($p = 0.0177$), while allicin and FB$_1$+allicin resulted in 1.24-fold and 1.49-fold decreases respectively relative to control (11.14 ± 0.7021).
Figure 4.5B: Decreased levels of RNS formation in all three HEK293 cell treatments relative to the control (11.14 ± 0.7021).

4.6 Thiobarbituric Acid Reactive Substances (TBARS) Assay

ROS-induced lipid peroxidation as an oxidative stress marker was assessed by quantifying extracellular levels of MDA. Extracellular MDA levels were significantly reduced for all treatments of Hek293 cells (Figure 4.6). A 1.92-fold decrease in FB₁ (p < 0.0001), 2.28-fold decrease in allicin (p < 0.0001) and a 1.27-fold decrease in combined FB₁ + allicin (p = 0.0001) treated cells was noted compared to the control (0.5855 ± 0.009887).
Figure 4.6: Significantly reduced levels of extracellular MDA in all HEK293 treatments relative the control (0.5855 ± 0.009887) - *** significantly different from the control.

4.7 Glutathione (GSH) Assay

The GSH concentration was measured as a marker for intracellular antioxidant capacity. The results showed that FB₁ did not significantly alter GSH concentration (Figure 4.7). However, a 1.34-fold decrease and 1.66-fold decrease in the allicin ($p = 0.0017$) and combined FB₁+allicin ($p = 0.0025$) treated cells was noted compared to the control (1.225 ± 0.02307) (Figure 4.7).
Figure 4.7: GSH concentrations relative to the control (1.225 ± 0.02307) show slight decrease in FB1 treatment and significant decrease in both allicin (p = 0.0017) and combined FB1+allicin (p = 0.0025) treated HEK293 cells.

4.8 Single Cell Gel Electrophoresis (SCGE) Assay

The comet assay was used to measure the length of comet tails as an indicator of DNA damage. All HEK293 cell treatments increased DNA damage shown by the significant increase in comet tail length (Figure 4.8). FB1 increased by 1.5-fold (p < 0.0001), allicin increased by 1.29-fold and FB1+allicin increased by 1.47-fold compared to the control (8.634 ± 0.4910).
Figure 4.8: Control cells (A) displayed an intact core of DNA, while FB$_1$ (B), allicin (C) and combined treatment (D) all caused migration of fragmented DNA from the nucleus, forming a comet tail. Significantly increased comet tail lengths were visualised for all HEK293 treatments relative to the control (8.634 ± 0.4910).
4.9 Caspases

The activity of caspases 8, 9 and 3/7 were assessed using luminometry. Caspases 8 (Figure 4.9A) and 9 (Figure 4.9B) demonstrate a significant upregulation in all Hek293 cell treatments: caspase 8 increased by 1.68-fold in FB1 (p = 0.0003), 1.94-fold in allicin (p = 0.0008) and 2.33 in FB1+allicin (p = 0.0023) treated cells relative to the control (2204000 ± 10750), and caspase 9 increased by 1.53-fold in FB1 (p < 0.0001), 1.85-fold in allicin (p < 0.0001) and 2.24-fold in FB1+allicin treated cells (p < 0.0001) relative to the control (5112000 ± 56180). Caspase 3/7 activity (Figure 4.9C) was significantly increased by 2.19-fold in the allicin-treated cells (p < 0.0001) and doubled in the combined treatment FB1 + allicin treatments (p < 0.0001) when compared to the control (261700 ± 2835).
Figure 4.9: Caspase activity following cell exposure to FB1, allicin and FB1+allicin. (C) Executioner caspase 3/7 showed a slight decrease in FB1 treated cells and a significant increase in subsequent treatments ($**p < 0.0001$). (B) Initiator caspase 8 (A) and 9 were significantly upregulated for all treatments.
4.10 Western blots

Western blotting was used to determine oxidative stress (Figure 4.10A) and apoptotic protein (Figure 4.10B) expression.

Oxidative stress markers included SOD2, catalase, GPx1, Nrf2 and HSP70 (Figure 4.10A). SOD2 was significantly downregulated for all Hek293 cell treatments presenting with a 1.42-fold decrease in FB1, a 1.72-fold decrease in allicin and a 2.75-fold decrease in FB1+allicin relative to the control (2.208 ± 0.03848). Likewise, catalase expression was decreased for all treatments, and the most significant depletion was noted in the combined treatment. In contrast, all three treatments significantly upregulated the expression of GPx1 relative to the control, with the upregulation less significant in the combined treatment. The expression of Nrf2 was also upregulated with a synergistic effect in the FB1+allicin compared to the control (0.3734 ± 0.1084). HSP70 expression was not markedly deviated from the control, with only slight increases occurring in the FB1 and combined treatments.

Apoptotic markers probed for were Bax, cPARP and p53 (Figure 4.10B). Bax was significantly increased relative to the control. Allicin increased Bax expression slightly more than FB1 alone. However, allicin treatment did not result in PARP cleavage, as was noted in the FB1 and FB1+allicin treatments. p53 was increased by 1.22-fold for FB1, but the allicin increase was not significant and the combined FB1+allicin treatment was unchanged when compared to the control.
Figure 4.10A: Protein expression for oxidative stress following HEK293 treatments. Nrf2 was upregulated for all treatments relative to the control (0.3734 ± 0.1084). HSP70 was only decreased in the allicin treatment. The expression of GPx1 was upregulated in all treatments. SOD2 expression was significantly downregulated. Catalase was overall decreased for all treatments.
Figure 4.10B: Protein expression for apoptotic markers following Hek293 cell treatments. BAX was significantly increased for all treatments. p53 was upregulated for both FB₁ and allicin at 1.22-fold and 1.06-fold respectively. cPARP expression was increased for all treatments relative to the control (0.4867 ± 0.02153).

4.11 Quantitative- Polymerase Chain Reactions (qPCR)
Hek293 treatments were assessed to determine the mRNA levels of SOD2 (Figure 4.11). FB₁ and allicin increased mRNA levels. The combined FB₁+allicin treatment decreased the mRNA levels of SOD2 relative to the control (0.9725 ± 0.02496).
Figure 4.11: SOD2 was increased for both FB₁ and allicin at 1.02-fold and 5.87-fold respectively. The combined FB₁+allicin treatment was significantly decreased.
FB₁ is a vastly studied mycotoxin produced as a secondary metabolite of *Fusarium verticillioides* known to contaminate crops. Studies have shown FB₁ to be toxic in multiple cell lines of human and animal origin (Berek *et al.*, 2001, Schmale and Munkvold, 2009). Although its mechanism of action remains unclear, many investigations performed on animal models demonstrate its effects on sphingolipid metabolism (Riley *et al.*, 1994) and more recently, a new epigenetic influence (Chuturgoon *et al.*, 2014) has been suggested. Overall, these mechanisms may result in dysregulation of protein synthesis and a variety of cytotoxic effects that include oxidative stress and cell death. The ubiquitous mycotoxin, FB₁, is geographically associated with OC in humans and liver/kidney cancer in rats (Voss *et al.*, 1993, Howard *et al.*, 2001, Khan *et al.*, 2018b). A recent study investigated its effects in HepG2 liver cancer cells (Chuturgoon *et al.*, 2014). However, its role in kidney toxicity in humans has not been investigated.

Garlic, a common medicinal plant has been used to treat various ailments over centuries. Its biological component allicin has been found to be responsible for many health benefits including anti-cancer properties through inhibition of cellular proliferation, inducing apoptosis and antioxidant activity to alleviate oxidative stress (Mikaili *et al.*, 2013, Gruhlke *et al.*, 2016). This study therefore sought to investigate the toxic effects of FB₁ in Hek293 cells. In addition, the ability of allicin to ameliorate their effects was studied.

Cytotoxicity was investigated using the MTT assay, which yielded an IC₅₀ of 215 µM and 3.905 µM for FB₁ and allicin respectively (Figure 4.1 A and B). The IC₅₀ in other studies varied for different cell lines, eliciting effects at different concentrations (Agarwal, 1996, Ha and Yuan, 2004, Liao *et al.*, 2009, Suddek, 2014) The decreased cell viability indicates that both FB₁ and allicin negatively influenced the mitochondrial ability to oxidise the MTT salt to its formazan product. Various studies
have shown that at varying concentrations, FB₁ decreases viability (Rumora et al., 2002, Chuturgoon et al., 2015). However, Myburg et al. (2002) showed that FB₁ had limited cytotoxicity in human OC SNO cells over 24 h and 48 h respectively (Myburg et al., 2002). With regards to allicin, studies have shown that it inhibits cell proliferation. According to Prager-Khoutorsky et al. (2007) this may be due to its effects on tubulin, thus the mitotic spindle will not form and cell division will be inhibited (Prager-Khoutorsky et al., 2007, Gruhlke et al., 2016).

Mitochondrial function is pivotal both in the production of ATP and activation of the intrinsic pathway of apoptosis. In addition, mitochondria also produce ROS as by-products of energy metabolism. Thus, determination of mitochondrial health is important. In this study, only allicin demonstrated increased mitochondrial integrity relative to the control (Figure 4.3). Furthermore, ATP production was increased by all treatments (Figure 4.4). The increased ATP implies an active electron transport chain. Where mitochondrial integrity is compromised, that will inevitably result in the production of ROS. Allicin increased mitochondrial integrity. When ATP production is increased in intact mitochondria, there would be less leakage of electrons and therefore less iROS formed (Brookes et al., 2004), as demonstrated by the results. iROS were increased by FB₁ and only slightly increased for allicin, while allicin failed to prevent iROS production in the presence of FB₁ (Figure 4.5A); in both instances, mitochondrial integrity was slightly lower relative to the control. However, RNS were decreased for all treatments (Figure 4.5B). Oxidative stress results from the production of ROS and an inadequate antioxidant response. In this study, the presence of oxidative stress is suggested by increased HSP70 in all treatments, where it appears that allicin potentiates the effects of FB₁, as well as Nrf2 which was synergistically increased in the combined treatment (Figure 4.10A).

An antioxidant response includes mobilisation of SOD2, catalase, GPx1, GSH and Nrf2. While downregulation of SOD2 protein expression occurred in all treatments (Figure 4.10A) its gene expression particularly increased (Figure 4.11) in the allicin treatments. Therefore, superoxide
conversion to H$_2$O$_2$ could not take place using SOD2. However, it may have been converted intracellularly by SOD1, a cytosolic enzyme isoform of SOD, into H$_2$O$_2$ (Birben et al., 2012). Khan et al. (2018b) demonstrated a decreased expression of SOD2 at 0-20 µM FB$_1$ doses in SNO cells (Khan et al., 2018a). In addition, underlying pathways including protein degradation play a major role in mRNA gene expression and protein concentration. This is supported by Scholze et al. (2011) whose study found an increase in the gene expression of SOD1 in chronic kidney disease patients coupled with decreased protein expression (Scholze et al., 2011, Krueger et al., 2016). Catalase, the antioxidant responsible for the conversion of H$_2$O$_2$ to H$_2$O was also decreased in the FB$_1$ and in the combined treatment.

The increased iROS levels therefore were not associated with SOD2 or catalase. The principle intracellular antioxidant is GSH; its depletion is usually an indication of oxidative stress (Forman et al., 2009). GSH was decreased by allicin and in the combined treatment (Figure 4.7), this is not surprising since allicin is a thiol agent that is known to oxidise GSH to GSSG, this has been confirmed by Gruhlke et al. (2016) who showed a decrease in GSH following administration of allicin (Gruhlke et al., 2016). In addition, GPx1 was increased (Figure 4.10A), which may account for the depletion of GSH; when GPx1 is activated to convert H$_2$O$_2$ to H$_2$O, GSH is oxidised to GSSG. The reduction of GSSG requires glutathione reductase (GR) and NADPH, a cofactor derived from the pentose phosphate pathway (Patra and Hay, 2014).

In the presence of cellular stress, HSP70 is expressed as a primary response sensor following protein misfolding (El Golli-Bennour and Bacha, 2011). This makes HSP70 a biomarker for oxidative stress. In this study, HSP70 indicated the presence of oxidative stress in the FB$_1$ and combined treatment (Figure 4.5A). While an increase in ROS production (H$_2$O$_2$) is noted in the FB$_1$ and combined treatment (Figure 4.5A), the antioxidant response is mobilised to ameliorate this effect. This is indicated by a depletion in the cellular antioxidants such as GSH, SOD and catalase (Figure 4.7, 10A).
Increased levels ROS are known to inactivate or deplete antioxidant enzymes (Lushchak, 2015), resulting in an accumulation or inadequate removal of ROS and oxidative stress (Fukai and Ushio-Fukai, 2011). For the allicin treatment, this utilisation of antioxidants kept H$_2$O$_2$ within normal limits, and oxidative stress was not present as indicated by HSP70 protein expression (Figure 4.10A). Increased Nrf2 (Figure 10A) explains the upregulation in the expression of GPx1 which in turn used up GSH to detoxify H$_2$O$_2$ (Ma, 2013), and is supported by research which shows that allicin elevates the levels of Nrf2 for the expression of other antioxidants to alleviate oxidative stress (Bat-Chen et al., 2010). A study on HepG2 cells demonstrated that transcription of antioxidant proteins was regulated by Nrf2 in FB$_1$-treated cells (García-Trejo et al., 2016, Arumugam et al., 2018). However, the Nrf2 response that results in increased SOD gene expression in allicin-treated cells, does not occur in the combined treatments (Figure 4.11). Therefore, despite the enhanced protein expression of Nrf2 following combination treatment, an accompanying replenishment of SOD will not occur. Thus allicin will not ameliorate the effects of FB$_1$. Given the reduced SOD gene expression relative to the FB$_1$-treatment suggests that allicin may increase oxidative stress in FB$_1$-treated cells.

While iROS were increased, there was no corresponding increase in lipid peroxidation (Figure 4.6). This decrease triggers the upregulation of antioxidant response molecules to maintain cellular homeostasis (Ayala et al., 2014). However, there was increased DNA fragmentation (Figure 4.8), which may be mediated by oxidative stress. When there is DNA damage, p53 is activated to initiate repair, cause cell cycle arrest or apoptosis. p53 was only slightly increased by FB$_1$ and allicin (Figure 4.10B) and was decreased in the combined treatments therefore repair mechanisms were not activated by allicin. DNA fragmentation is a morphological feature of apoptosis; therefore, in this study it may represent a late-stage marker for apoptosis.

Apoptosis is a programmed cell death mechanism facilitated by a family of proteins called caspases; in this study caspase activities were evaluated. In all treatments, the initiator caspases 8 and 9 were
upregulated (Figure 4.9A, B) which suggests that apoptosis is initiated via the extrinsic and intrinsic pathways; indeed, both FB\textsubscript{1} and allicin have been shown to induce apoptosis (Dragan \textit{et al}., 2001, Oommen \textit{et al}., 2004, Minervini \textit{et al}., 2004, Bat-Chen \textit{et al}., 2010) Upon activation, these caspases initiate the apoptotic cascade that culminates in the cleavage of caspase 3/7 known as the executioner of apoptosis. However, the results show that FB\textsubscript{1} did not increase caspase 3/7 (Figure 4.9C). It is possible that inhibitors of apoptosis (IAPs) may have blocked caspase 3/7 activation, by blocking apoptosis downstream of cytochrome c release by binding to and inhibiting active caspase 9 within the apoptosome, preventing downstream activation of procaspase 3/7. Chuturgoon \textit{et al}., (2015) showed that BIRC 8, an IAP, was increased following FB\textsubscript{1} administration in HepG2 cells, which may have been responsible for lack of execution caspase 3/7 activity despite caspase 9 activation (Chuturgoon \textit{et al}., 2015). In addition the presence of mitochondrial proteins Smac/DIABLO and Omi/HtrA2 antagonise this inhibitory action (Burke \textit{et al}., 2010, Parrish \textit{et al}., 2013). Furthermore, HSP70 is an anti-apoptotic protein that can inhibit apoptosis upstream of caspase 3/7 activation; this can therefore be supported by the observed increase in HSP70 in FB\textsubscript{1} treated cells (Figure 4.10A). A study on animal models also showed that FB\textsubscript{1} increased the expression of HSP70 and this is inclined with our findings (Kócsó \textit{et al}., 2018).

An overall increase in caspase levels was noted in allicin and combined allicin and FB\textsubscript{1}-treated Hek293 cells (Figure 4.9C) suggesting caspase-dependant activation of apoptosis. These findings are in agreement with a study done by Oommen \textit{et al}., (2004) on human cervical and colon cancer cells wherein allicin halted cancer growth through induction of caspase-dependent apoptosis. Their results showed an increase in 3, 8 and 9 as well as PARP cleavage (Oommen \textit{et al}., 2004). Bax is responsible for the formation of mitochondrial pores to release cytochrome c. Our results demonstrated an overall increase in Bax expression (Figure 4.10B). Alternatively, mitochondrial toxicity associated with allicin treatment could be responsible for cytochrome c release and caspase activation.
Activated caspase 3/7 results in the cleavage of iCAD and PARP forming CAD and cPARP respectively. CAD is a DNase that cleaves DNA into 180 bases per unit and is associated with the fragmentation of DNA as seen in Figure 4.9. The PARP are a family of proteins that assist in DNA damage and repair by binding to and repairing detected single and double-stranded DNA breaks. Caspase 3 cleaves PARP and therefore renders it inactive. Oommen et al. (2004) showed increased cPARP in human cervical and human colon cancer cells (Oommen et al., 2004). Likewise, 10 µM FB₁ treatment of SNO cells also caused an increase in cPARP (Khan et al., 2018a). Thus, allicin increased induction of apoptosis in the presence of FB₁, which was expected since oxidative stress was present. If increased apoptosis is accompanied by compensatory cell proliferation to replenish damaged cells, then cancer could result. This is of particular concern since the combined treatment also decreased DNA repair.
CHAPTER 6: CONCLUSION

This study sought to explore mechanisms of toxicity exerted by FB$_1$ in a human kidney cell line coupled with the potential therapeutic properties of allicin on FB$_1$ treated Hek293 cells. The results show that FB$_1$ and allicin cause a decrease in cell viability. All three treatments increased MDA in conjunction with increased ATP production. However, H$_2$O$_2$ concentration was not altered by allicin. The activation of an antioxidant response occurred; Nrf2 and GPx1 were upregulated but SOD2 and catalase were decreased, suggesting alternative modes of oxidant detoxification. Apoptosis was triggered in all treatments via both intrinsic and extrinsic pathways, but execution in the FB$_1$ treatment was not caspase-dependent. Cumulatively, FB$_1$ and allicin individually trigger apoptosis, but only FB$_1$ induced oxidative stress. When combined, a possible synergistic relationship is demonstrated in the expression of proteins. In addition, allicin caused decreased SOD2 gene expression in the presence of FB$_1$ despite increased Nrf2 upregulation. Therefore, allicin did not exert a therapeutic effect on FB$_1$-treated Hek293 cells.

Following findings from this study, further investigations on the effects of IAP in FB$_1$-induced apoptosis is recommended. The inflammatory pathway as a consequence of both allicin and FB$_1$ on $\textit{in vitro}$ and $\textit{in vivo}$ models is also of interest for future studies.

While more research is required, allicin should not be recommended as a supplement to ameliorate the cytotoxic effects of FB$_1$. This study only tested the IC$_{50}$ concentrations of FB$_1$ and allicin. Perhaps expanding the range of treatment concentrations can give a better picture of the interaction between FB$_1$ and allicin.
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APPENDICES

Appendix A: Nitric oxide assay

![NOS Assay Standard curve](image)

**Figure 1:** Mean absorbance values of NOS standard curve

Appendix B: Glutathione (GSH) Assay
Figure 1: Mean absorbance values of GSH concentrations standard curve.

Appendix C: Protein standardisation

Figure 1: Standard curve for protein standardisation