The effects of melatonin supplementation on vascular tissue during first line ART: an *in vivo, ex vivo* and *in vitro* study

by
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*Thesis presented in fulfilment of the requirements for the degree of Master of Science in the Faculty of Health Science at Stellenbosch University*

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Co-supervisor: Prof Hans Strijdom

March 2018
Declaration

By submitting this thesis electronically, I declare that the entirety of the work contained therein is my own, original work, that I am the sole author thereof (save to the extent explicitly otherwise stated), that reproduction and publication thereof by Stellenbosch University will not infringe any third party rights and that I have not previously in its entirety or in part submitted it for obtaining any qualification.

March 2018
Abstract

Introduction: Although Antiretroviral therapy (ART) has dramatically reduced HIV-associated morbidity and mortality, non-HIV-related conditions and comorbidities continue to rise in this population. Cardiovascular disease (CVD) has been reported to be the leading cause of death in the HIV-positive population receiving ART. ART is thought to impair vascular endothelial function through increased reactive oxygen species (ROS) and reactive nitrogen species (RNS) production. In this study, we aim to assess the effects of melatonin - a potent antioxidant -supplementation during ART on specific intracellular products of rat Aortic Endothelial Cells (AECs) as well as on the vascular reactivity of rat aortas.

Methods: Cells were serum starved and treated with three different melatonin (1nM; 1μM; 10μM) or ART (Low: EFV: 5μM; FTC: 5μM; TDF: 80nM; Mid: EFV: 8μM; FTC: 7.5μM; TDF: 400nM; High: EFV: 12μM; FTC: 10μM; TDF: 1μM) concentrations for up to 24 hours. Nitric oxide (NO), RNS and necrosis were measured with a platereader (control expressed as 100%). The concentration that resulted in the greatest differences, compared to untreated cells, was selected for co-treatment studies and protein investigations, where the same parameters were measured. The effects of acute melatonin and ART administration on vascular reactivity was measured by aortic ring isometric tension studies in aortas extracted from control male Wistar rats. The endothelium-dependent and independent vascular reactivity was measured by isometric tension studies on aortas harvested from male Wistar rats that were treated for 8 weeks with melatonin (10mg/kg/day) and/or ART (EFV: 51.6 mg/kg; FTC: 17.4 mg/kg; TDF: 25.8 mg/kg). Signalling proteins involved in these changes were measured by western blot analyses.

Results (Mean±SEM): Dose Response Studies: 1nM melatonin decreased necrosis [92.56±3.11%;p=0.0004] compared to untreated controls [100.00±1.04%; p=0.0004]. High-concentration ART lead to increased NO production [112.70±2.17%; Control:100.00±2.22%;p=0.0015], RNS production [108.80±2.20%; Control:100±0.79%;p=0.0152] and necrosis [107.30±1.34%; Control: 100.00±0.86%;p=0.0251], compared to untreated controls.

Main Studies: 1nM melatonin decreased necrosis [92.43±3.75%;p=0.0002], while High-concentration ART increased necrosis [121.30±9.11%;p=0.0002] compared to untreated cells [100.00±1.07%;p=0.0002]. When combined, 1nM melatonin + High-concentration ART decreased necrosis [94.17±5.08%;p=0.0002] compared to ART alone.
Western blot analyses (arbitary units) showed that ART increased nitrotyrosine levels [2.31±0.34; Control:1.00±0.18;p=0.0486], but decreased p22 PHOX [0.20±0.043; Control:1.00±0.15;p<0.0001], and cleaved caspase-3 [0.25±0.038; Control:1.00±0.18;p=0.0005], expression.

In acute aortic ring experiments, ART exposure elicited a burst of contraction during the treatment period, followed by a significant attenuation in accumulative contraction compared to all other groups. In endothelium-dependent and independent contraction studies on aortas from treated rats, all groups showed a pro-contractile response compared to the control. Western blot analyses showed that ART decreased cleaved caspase-3 [0.27±0.08;p=0.0055] expression.

**Conclusion:** Decreased necrosis in AECs treated with combined melatonin and ART, compared to AECs treated with ART alone shows the protective effect of melatonin. Further specific protein investigations are needed to elucidate this mechanism. Western blots showed that ART induced anti-apoptotic effects and increased RNS production, but not NADPH-oxidase activity. The initial contractile burst following acute ART exposure may precondition the aortas, resulting in the decreased accumulative contractile capacity. Chronic ART treatment studies showed that ART treatment does not seem to affect vasorelaxation. Blot data reconfirmed that ART is also anti-apoptotic *in vivo.*
Abstrak

**Inleiding:** Alhoewel antiretrovirale terapie HIV verwante morbiditeit en mortaliteit dramaties verminder het, styg nie-HIV-verwante toestande in hierdie populasie. Kardiovaskulêre siektes word beskou as die hoof oorsaak van sterftes in die HIV positiewe populasie wat antiretrovirale terapie (ART) ontvang. ART kan die vaskulêre endoteelfunksie benadeel deur verhoogde reaktiewe suurstof species (ROS) en reaktiewe stikstof species (RNS) produksie. In hierdie studie was die effekte van melatonien (kragtige antioksidant) aanvulling gedurende ART ondersoek, met spesifieke verwysing na intrasellulêre afskeidingsprodukte van rot aorta endoteelselle, asook in die konteks van rot aorta vaskulêre reaktiwiteit.

**Metodes:** Selle was onderwerp aan serum weerhouding en behandel met drie verskillende melatonien (1nM; 1uM; 10uM) of ART (**Laag**: EFV: 5µM; FTC: 5µM; TDF: 80nM; **Medium**: EFV: 8µM; FTC: 7.5µM; TDF: 400nM; **Hoog**: EFV: 12µM; FTC: 10µM; TDF: 1µM) konsentrasies vir tot 24uur. Stikstofoksied (NO), RNS and nekrose is gemeeet met 'n plaatleser (kontrole uitgedruk as 100%). Die konsentrasie wat gelei het tot die grootste verskille, in vergelyking met die onbehandelde selle, was gekies vir ko-behandeling studies en proteïen ontledings, waar dieselfde veranderlikes gemeet was. Die effekte van akute melatonien en ART toediening op vaskulêre reaktiviteit is gemee d.m.v. aortiese ring isometriese spanningstudies in aortas afkomstig van gesonde manlike Wistar rotte. Die endoteel-afhanklike en onafhanklike vaskulêre reaktiwiteit is gemee d.m.v. isometriese spanningstudies in aortas afkomstig van manlike Wistar rotte wat vir 8 weke behandel was met melatonien (10mg/kg/day) en/of ART (EFV: 51.6 mg/kg; FTC: 17.4 mg/kg; TDF: 25.8 mg/kg). Seinproteïne in hierdie weefsel is gemeet deur Western blot analises.

**Resultate (Mean±SEM):** Dosis-reaksie Ondersoekte: 1nM melatonien het nekrose [92.56±3.11%; p=0.0004] laat verminder in vergelyking met onbehandelde kontrole [100.00±1.04%; p=0.0004]. Hoë-konsentrasie ART het tot verhoogde NO produksie [112.70±2.17%; kontrole:100.00±2.22%; p=0.0015], RNS produksie [108.80±2.20%; kontrole:100.00±0.79%; p=0.0152] en nekrose [107.30±1.34%; kontrole: 100.00±0.86%; p=0.0251] gelei, in vergelyking met onbehandelde kontrole.

Hoof Ondersoekte: 1nM melatonien nekrose [92.43±3.75%; p=0.0002] laat toeneem, terwyl hoë-konsentrasie ART nekrose [121.30±9.11%; p=0.0002] in vergelyking met onbehandelde selle [100.00±1.07%; p=0.0002] verhoog. Tydens gekombineerde
behandeling, het 1nM melatonien + Hoë-konsentrasie ART nekrose [94.17±5.08%; p=0.0002] verlaag teenoor ART alleen. Western blot analyses (arbitreëre eenhede) het getoon dat ART nitrotirosien vlakke vermeerder het [2.31±0.34; kontrole:1.000±0.18;p=0.0486], maar p22 PHOX [0.20±0.04; kontrole:1.00±0.15;p<0.0001], en gekliefde kaspase-3 [0.25±0.038; kontrole:1.00±0.18;p=0.0005] uitdrukking verlaag het. In akute aorta ring eksperimente, het ART blootstelling ‘n skielike en kortstondige kontraksie gedurende die behandelings periode veroorsaak, gevolg deur ‘n beduidende verswakking in akkumulatiewe kontraksie teenoor al die ander groepe. In endoteel-afhanklike en onafhanklike kontraksie studies op aortas van behandelde rotte, het alle groepe ‘n pro-kontraktiele reaksie getoon in vergelyking met die kontrole. Western blot analyses het getoon dat ART v gekliefde kaspase-3 uitdrukking verlaag het [0.27±0.08;p=0.0055].

**Gevolgtrekking:** Die verlaagde nekrose wat in aorta endoteelselle (AECs), behandel met gekombineerde melatonien en ART, waargeneem is teenoor AECs wat slegs ART ontvang het, is ‘n bewys van die beskermende effekte van melatonien. Toekomstige studies wat spesifieke proteïene ondersoek is nodig om die mechanisme verder te verklar. Western blots ondersoek het gesien dat ART anti-apoptotiese effekte en vermeerder RNS produksie veroorsaak, maar geen effekte op NADPH-oksidase aktiwiteit toon nie. Die aanvanklike kontraktiele respons tydens akute ART blootstelling kan die aortas vooraf kondisioneer wat ‘n verlaagde akkumulatiewe kontraktiele kapasiteit veroorsaak het. Chroniese ART behandeling studies het getoon dat die behandeling nie die bloedvatverslapping beïnvloed nie. Die Western blot resultate van die in vivo studies het die anti-apoptotiese effekte van ART herbevestig.
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<tr>
<td>+C</td>
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<td>L-arginine</td>
</tr>
<tr>
<td>L-Cit</td>
<td>L-citrulline</td>
</tr>
<tr>
<td>L-NAME</td>
<td>Nω-Nitro-L-arginine methyl ester</td>
</tr>
<tr>
<td>LOX</td>
<td>Lipooxygenases</td>
</tr>
<tr>
<td>M</td>
<td>Muscarinic receptor</td>
</tr>
<tr>
<td>MeOH</td>
<td>Methanol</td>
</tr>
<tr>
<td>MI</td>
<td>Myocardial infarction</td>
</tr>
<tr>
<td>MLCK</td>
<td>Myosin light chain kinase</td>
</tr>
<tr>
<td>Abbr.</td>
<td>Full Form</td>
</tr>
<tr>
<td>-------</td>
<td>-----------</td>
</tr>
<tr>
<td>mRNA</td>
<td>Messenger-ribonucleic acid</td>
</tr>
<tr>
<td>mtPTP</td>
<td>Mitochondrial permeability transition pore</td>
</tr>
<tr>
<td>NADPH</td>
<td>Nicotinamide adenine dinucleotide phosphate</td>
</tr>
<tr>
<td>NFV</td>
<td>Nelfinavir</td>
</tr>
<tr>
<td>nNOS</td>
<td>Neuronal Nitric Oxide Synthase</td>
</tr>
<tr>
<td>NNRTIs</td>
<td>Non-Nucleoside reverse transcriptase inhibitors</td>
</tr>
<tr>
<td>NO</td>
<td>Nitric oxide</td>
</tr>
<tr>
<td>NO(_2^-)</td>
<td>Nitrite</td>
</tr>
<tr>
<td>NOS</td>
<td>Nitric oxide synthase</td>
</tr>
<tr>
<td>NOX</td>
<td>Nicotinamide adenine dinucleotide phosphate oxidase</td>
</tr>
<tr>
<td>NRTIs</td>
<td>Nucleoside reverse transcriptase inhibitors</td>
</tr>
<tr>
<td>NtRTIs</td>
<td>Nucleotide reverse transcriptase inhibitors</td>
</tr>
<tr>
<td>NVP</td>
<td>Nevirapine</td>
</tr>
<tr>
<td>O(_2)</td>
<td>Oxygen</td>
</tr>
<tr>
<td>O(_2^-)</td>
<td>Superoxide anion</td>
</tr>
<tr>
<td>OH(^-)</td>
<td>Hydroxyl</td>
</tr>
<tr>
<td>ONOO(^-)</td>
<td>Peroxynitrite</td>
</tr>
<tr>
<td>ONOOCO(_2^-)</td>
<td>Nitroso-peroxocarboxylate</td>
</tr>
<tr>
<td>ONOOH</td>
<td>Peroxynitrous acid</td>
</tr>
<tr>
<td>ORAC</td>
<td>Oxygen radical absorbance capacity</td>
</tr>
<tr>
<td>PAOD</td>
<td>Peripheral arterial occlusive disease</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate-buffered saline</td>
</tr>
<tr>
<td>PGH(_2)</td>
<td>Prostaglandin H2</td>
</tr>
<tr>
<td>PGI(_2)</td>
<td>Prostaglandin I2</td>
</tr>
<tr>
<td>Phe</td>
<td>Phenylephrine</td>
</tr>
<tr>
<td>PI</td>
<td>Propidium iodide</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>PIs</td>
<td>Protease inhibitors</td>
</tr>
<tr>
<td>PKA</td>
<td>Protein kinase A</td>
</tr>
<tr>
<td>PKC</td>
<td>Protein kinase C</td>
</tr>
<tr>
<td>R3-IGF-1</td>
<td>Long chain human insulin-like growth factor</td>
</tr>
<tr>
<td>rhEGF</td>
<td>Recombinant human epidermal growth factor</td>
</tr>
<tr>
<td>rhFGF-B</td>
<td>Recombinant human fibroblastic growth factor B</td>
</tr>
<tr>
<td>rLPV</td>
<td>Lopinavir/ritonavir</td>
</tr>
<tr>
<td>RNS</td>
<td>Reactive nitrogen species</td>
</tr>
<tr>
<td>ROS</td>
<td>Reactive oxygen species</td>
</tr>
<tr>
<td>RPM</td>
<td>Revolutions per minute</td>
</tr>
<tr>
<td>RT</td>
<td>Reverse transcriptase</td>
</tr>
<tr>
<td>RTV</td>
<td>Ritonavir</td>
</tr>
<tr>
<td>S1B</td>
<td>Serotonin receptor</td>
</tr>
<tr>
<td>SA</td>
<td>South Africa</td>
</tr>
<tr>
<td>SABS</td>
<td>South African Bureau of Standards</td>
</tr>
<tr>
<td>SDS</td>
<td>Sodium dodecyl sulphate</td>
</tr>
<tr>
<td>SEM</td>
<td>Standard error of the mean</td>
</tr>
<tr>
<td>sGC</td>
<td>Soluble guanylyl cyclase</td>
</tr>
<tr>
<td>SMART</td>
<td>The Strategies for Management of Anti-Retroviral Therapy</td>
</tr>
<tr>
<td>SNP</td>
<td>Sodium nitroprusside</td>
</tr>
<tr>
<td>SOCa²⁺</td>
<td>Store-operated Ca²⁺ channel</td>
</tr>
<tr>
<td>SOD</td>
<td>Superoxide dismutases</td>
</tr>
<tr>
<td>SP</td>
<td>Substance P</td>
</tr>
<tr>
<td>SQV</td>
<td>Saquinavir</td>
</tr>
<tr>
<td>SR</td>
<td>Sarcoplasmic reticulum</td>
</tr>
<tr>
<td>SS</td>
<td>Serum starved</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------------------------------------</td>
</tr>
<tr>
<td>Tat</td>
<td>Trans-activator of transcription</td>
</tr>
<tr>
<td>TDF</td>
<td>Tenofovir disoproxil fumarate</td>
</tr>
<tr>
<td>TP</td>
<td>Thromboxane prostanoid receptor</td>
</tr>
<tr>
<td>TXA₂</td>
<td>Thromboxane A</td>
</tr>
<tr>
<td>UCT</td>
<td>University of Cape Town</td>
</tr>
<tr>
<td>US</td>
<td>University of Stellenbosch</td>
</tr>
<tr>
<td>USA</td>
<td>United States of America</td>
</tr>
<tr>
<td>VEGF</td>
<td>Vascular endothelial growth factor</td>
</tr>
<tr>
<td>VSMC</td>
<td>Vascular smooth muscle cells</td>
</tr>
</tbody>
</table>
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1 - Literature Review

1.1 General Introduction

The human immunodeficiency virus (HIV) is a global pandemic affecting 34 million people, with African countries accounting for 70% of all HIV-infected individuals worldwide (Thienemann, Sliwa, & Rockstroh, 2013). In South Africa (SA) alone, there are approximately 7 million people living with HIV (Statistics South Africa, 2015). There is still no cure for HIV infection, however, implementation of antiretroviral therapy (ART) has allowed for the effective management of the virus. Although ART has dramatically reduced HIV-associated morbidity and mortality, non-HIV-related conditions and comorbidities continue to rise (Thienemann et al., 2013). Cardiovascular disease (CVD) has been reported to be the leading comorbidity and cause of death in the HIV-positive population receiving ART (Thienemann et al., 2013), creating a double burden of disease – both communicable and non-communicable – specific to low-and-middle-income countries like SA.

Endothelial dysfunction (ED) is defined as an imbalance between vasodilating and vasoconstricting substances produced by, or acting on the endothelium (Deanfield et al., 2005), and is recognised as a critical initiating factor in CVD. Numerous studies on the effects of ART on the vasculature indicate that ART may impair the function of the vascular endothelium (Jiang et al., 2010). Increased production of reactive oxygen species (ROS) and imbalances between the levels of oxidants and antioxidants in the vasculature are considered important factors in ART-induced ED.

There is need for a therapy with the potential to alleviate the oxidative stress induced by ART. Melatonin is a free radical scavenger and strong antioxidant (Reiter, Tan, Osuna, & Gitto, 2000), giving it the potential to ameliorate damage caused by ART-induced ROS production, which leads to ED.

1.2 The Vascular Endothelium

The vascular endothelium is made up of a monolayer of endothelial cells (ECs), which line the lumen of blood vessels, providing a barrier between the vascular walls and circulating blood (Lerman & Zeiher, 2005; M. S. Park, Ravi, & Araujo, 2010). For many years, the endothelium was thought of as an inert membrane, with the sole purpose of maintaining
vessel wall permeability through the physical separation of the tissue and blood (Cines et al., 1998). Now, it is well established that the endothelium is in fact a dynamic organ that is heterogeneous and involved in the synthesis and secretion of key factors, as well as fulfilling important metabolic and immunological functions (Fishman, 1982). The endothelium plays a vital role in preserving vascular homeostasis by maintaining a delicate balance between vasodilatory and vasoconstrictory factors (Bevilacqua, Nelson, Mannori, & Cecconi, 1994; Versari, Daghini, Virdis, Ghiadoni, & Taddei, 2009). For this reason, maintaining endothelial health is fundamental in sustaining an antithrombotic and antiatherogenic milieu. Table 1.1 summarises the various atheroprotective effects of the healthy endothelium.

Table 1.1: A list of favourable effects enforced by a healthy endothelium (Bonetti, Lerman, & Lerman, 2003).

<table>
<thead>
<tr>
<th>Favourable and Atheroprotective Effects of the Healthy Endothelium:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Promotion of vasodilation</td>
</tr>
<tr>
<td>Antioxidant effects</td>
</tr>
<tr>
<td>Anti-inflammatory effects</td>
</tr>
<tr>
<td>Inhibition of leukocyte adhesion and migration</td>
</tr>
<tr>
<td>Inhibition of smooth muscle cell proliferation and migration</td>
</tr>
<tr>
<td>Inhibition of platelet aggregation and adhesion</td>
</tr>
<tr>
<td>Anticoagulant effects</td>
</tr>
<tr>
<td>Profibrinolytic effects</td>
</tr>
</tbody>
</table>

1.2.1 Localisation and Structure

The vascular endothelium lines the entire circulatory system, from the heart, to the smallest capillaries (Rajendran et al., 2013). Positioned at the interface between the blood and the tissue, the endothelium is in the ideal location to detect and respond to any biological changes in the local environment caused by trauma or inflammation (Cines et al., 1998). The vascular wall is comprised of three layers (Fig 1.1): an inner layer called the intima, a middle layer known as the tunica media, and an outer layer called the tunica adventitia (Sandoo, van Zanten, Metsios, Carroll, & Kitas, 2010). The endothelium
comprises approximately $1.6 \times 10^{13}$ cells, all of which are in close association with the smooth muscle cells (Limaye & Vadas, 2007), and located on the intima. These cells are highly specialised, with the ability to receive and transmit biochemical and physical information (Cines et al., 1998). This information can be transmitted directly – through active transport or direct permeation through the intercellular spaces between the ECs – or indirectly, whereby the ECs modulate the behaviour of the smooth muscle cells and other components of the vessel wall (Cines et al., 1998).

![Diagram of a healthy artery](image)

**Figure 1.1:** Morphologically distinct layers in a healthy artery (Lusis, 2000).

These actions are facilitated via membrane-bound receptors; which convey the signals of numerous molecules, such as proteins, lipid transporting particles, metabolites and hormones (Cines et al., 1998). The ECs also have specialised junctional proteins and receptors, which govern cell-cell and cell-matrix interactions.

### 1.2.2 Function

It has been established that the endothelium embodies a wide range of homeostatic functions (Durand & Gutterman, 2013). These functions are carried out through the secretion of specific mediators (Table 1.2) that have the ability to influence vascular hemodynamics (Cines et al., 1998). The various mediators controlled by ECs play an important role in the regulation of vascular tone and growth of blood vessels, thrombosis and thrombolysis, as well as platelet and leukocyte interactions with the vessel wall (Verhamme & Hoylaerts, 2006). ECs also contribute to the regulation of blood pressure and blood flow, by releasing vasodilators and vasoconstrictors (Cines et al., 1998).
**Table 1.2: Endothelium Derived Vaso-regulatory Mediators (Adapted from Cines et al., 1998; Mudau, Genis, Lochner, & Strijdom, 2012).**

<table>
<thead>
<tr>
<th>Mediator</th>
<th>Effects</th>
<th>Source and Mechanism of Action</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitric Oxide (NO)</td>
<td>- Potent vasodilator</td>
<td>• Synthesises by enzymes: eNOS; nNOS and iNOS, with eNOS the major endothelial source of NO during physiological conditions</td>
</tr>
<tr>
<td></td>
<td>- Inhibits inflammation, VSMC proliferation and migration, platelet activation, aggregation and adhesion, and leukocyte adhesion</td>
<td>• Diffuses from ECs to underlying VSMCs where it binds to sGC, leading to a cascade of events that ultimately result in vascular relaxation</td>
</tr>
<tr>
<td></td>
<td>- Regulates myocardial contractility</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Regulates cardiac metabolism</td>
<td></td>
</tr>
<tr>
<td></td>
<td>- Maintains basal tone of vessels</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td><strong>Nitric Oxide (NO)</strong></td>
</tr>
<tr>
<td>Prostacyclin (PGI2)</td>
<td>- Vasodilator</td>
<td>• Derived from arachidonic acid by cyclooxygenase-2 (COX-2)</td>
</tr>
<tr>
<td></td>
<td>- Inhibits platelet aggregation and deposition</td>
<td></td>
</tr>
<tr>
<td>Endothelium-derived hyperpolarising factor (EDHF)</td>
<td>- Vasodilator, particularly in small arteries (≤ 300 μm)</td>
<td>• Its identity is still under suspicion with proposed candidates such as potassium ions and hydrogen peroxide</td>
</tr>
<tr>
<td>Endothelin-1 (ET-1)</td>
<td>- Potent vasoconstrictor</td>
<td>• Synthesised by endothelin-converting enzyme</td>
</tr>
<tr>
<td></td>
<td>- Mitogen for VSMCs</td>
<td>• Exerts its effects via two receptors: ET&lt;sub&gt;A&lt;/sub&gt; expressed on endothelial cells which promote vasoconstriction and ET&lt;sub&gt;B&lt;/sub&gt; on VSMCs, which promote NO production and ultimately reduction in ET-1 production</td>
</tr>
<tr>
<td>Thromboxane A (TXA2)</td>
<td>- Potent vasoconstrictor</td>
<td>• Derived from arachidonic acid by COX-1</td>
</tr>
<tr>
<td>Angiotensin II (Ang II)</td>
<td>- Potent vasoconstrictor</td>
<td>• Synthesised by angiotensin converting enzyme</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Elicits its effects via two receptors: AT&lt;sub&gt;1&lt;/sub&gt; which promotes vasoconstriction and cell proliferation, and AT&lt;sub&gt;2&lt;/sub&gt; which antagonises the effects of AT&lt;sub&gt;1&lt;/sub&gt;</td>
</tr>
</tbody>
</table>

VSMC (vascular smooth muscle cell); eNOS (endothelial nitric oxide synthase); nNOS (neuronal nitric oxide synthase); iNOS (inducible nitric oxide synthase); sGC (soluble guanylyl cyclase).
1.2.2.1 Nitric Oxide and Nitric Oxide Synthase

As previously mentioned on page 3, there are numerous vasoactive substances produced by the ECs. Nitric Oxide (NO) is the most potent endothelium-dependant vasodilator synthesised in the endothelial cell layer (Durand & Gutterman, 2013; Mudau et al., 2012), and causes the relaxation of the underlying vascular smooth muscle cells (VSMCs) (Sandoo et al., 2010). NO is the heterodiatomic free radical product that is formed through the oxidation of the amino acid L-arginine, to L-citrulline (Fig.1.2) (K. Park & Park, 2015; Stamler, Singel, & Loscalzo, 1992). This conversion is facilitated by the enzyme nitric oxide synthase (NOS) (Palmer, Ashton, & Moncada, 1988).

Figure 1.2: The formation of NO from L-Arginine, with the NADPH and oxygen requirements for each reaction shown (Stuehr, 2004). NADPH (Nicotinamide adenine dinucleotide phosphate).

NOS exists in three different isoforms: neuronal NOS (nNOS); inducible NOS (iNOS) and endothelial NOS (eNOS). nNOS produces NO, which plays the role of a neuronal messenger that has the ability to regulate synaptic neurotransmitter release (Prast & Philippu, 2001), while iNOS is only expressed in injured cells that have been exposed to inflammatory mediators that can activate macrophages (Michel & Feron, 1997). eNOS is the most abundant isoform of NOS and is responsible for the NO produced in the vasculature, and therefore vessel dilation (Sandoo et al., 2010). The various isoforms of NOS were originally named and classified according to the cells they were initially observed in, however, it is now evident that the different isoforms also occur in numerous other cells, including cardiac myocytes (Balligand et al., 1995); as well as blood platelets, the hippocampus and skeletal muscle (Arnal, Dinh-Xuan, Pueyo, Darblade, & Rami, 1999).
In the vascular endothelium, NO production is influenced by the release of chemical agonists acting on specific endothelial chemo-receptors (K. Park & Park, 2015). Such NO agonists include acetylcholine (ACh), bradykinin (BK), adenosine tri-phosphate (ATP), adenosine di-phosphate (ADP), substance P (SP) and thrombin (Moncada & Higgs, 2006). Inactive eNOS is bound to the protein caveolin and kept within small invaginations in the cell membrane, known as caveolae (Bucci et al., 2000). eNOS is activated by the receptor-dependent NO agonists, which stimulate the release of calcium from the endoplasmic reticulum (ER) (Fig 1.4) (Bae et al., 2003), thereby increasing intracellular calcium levels and causing the detachment of eNOS from caveolin (Fig 1.3) (Sandoo et al., 2010; Venema, Sayegh, Arnal, & Harrison, 1995). When intracellular calcium from the ER is depleted, endothelial membrane receptors receive a signal to open the calcium channels and allow the entry of extracellular calcium into the cell (Schilling, Cabello, & Rajan, 1992), which is known as store operated calcium entry or capacitative calcium entry (Putney, 1986). In order to bind to and activate eNOS, calcium must undergo a structural change, which is achieved through the binding of calcium to calmodulin within the cell cytoplasm (Fleming & Busse, 1999). Active eNOS is then able to catalyse the production of NO.

Figure 1.3: eNOS signaling in caveolae. Depicted in the figure is the cycle of eNOS activation and inhibition in caveolae. In step 1, Cav-1 is bound to eNOS, inhibiting eNOS activity. In step 2, agonist activation by Ach initiates an influx of calcium that binds to and activates calmodulin. In step 3, calcium-activated calmodulin binds to eNOS thereby relieving its inhibition by Cav-1 and NO is produced. Finally, in step 4, Cav-1 binds to eNOS again, completing the cycle (Razani, Woodman, & Lisanti, 2002). Cav-1 (caveolin); Ca^{2+} (calcium).
of NO through the conversion of L-arginine (Fig 1.2). This process of NO production in ECs is shown in Figure 1.4.

In this particular pathway, it is important to note that NO production is dependent on the amount of intracellular calcium in the ER, as well as the quantity of extracellular calcium that diffuses into the cell (Sandoo et al., 2010). When calcium levels decrease, the calcium-calmodulin complex dissociates from eNOS. The unbound eNOS is then free to bind with caveolin again, and therefore becomes inactivated (Fleming & Busse, 1999). In this way, short-term increases in NO are dependent on calcium levels, but once intracellular calcium is depleted, alternative mechanisms need to be activated in order to further regulate NO production. In such cases, eNOS can be alternatively activated through phosphorylation (Butt et al., 2000), which occurs through specific protein kinases, like protein kinase A (PKA) (Bae et al., 2003) and cyclic guanosine-3’, 5-monophosphate (cGMP) protein kinase dependent II (Butt et al., 2000).

Once eNOS has been activated and NO has been consequently synthesised, NO diffuses across the EC into the adjacent VSMC (Fig 1.4), where it binds to the enzyme soluble guanylyl cyclase (sGC), thereby activating it (Ignarro, Harbison, Wood, & Kadowitz, 1986). Active sGC increases the conversion rate of guanosine triphosphate (GTP) to cGMP, which decreases smooth muscle tension (Jones, Wong, Jankowski, Akao, & Warner, 1999), effectively decreasing the release of calcium from the sarcoplasmic reticulum (SR) in the VSMCs (Collins, Griffith, Henderson, & Lewis, 1986), which overall reduces VSMC contraction. All the mechanisms described above are continuously active, and the resultant production of NO is used to maintain basal vasodilatory tone (Sandoo et al., 2010).
Under normal physiological conditions, endothelial NO production is mainly regulated by mechanical forces, particularly shear stress, which act on specific mechano-receptors (Gutiérrez et al., 2013). This mechanical activation of eNOS is the most important regulator of eNOS activity and contributes to the phenomenon of flow-mediated vasodilatation (Fig 1.4), which is a vital autoregulatory mechanism that increases blood flow in response to physical activity or exercise (Loscalzo & Vita, 1994).

NO production and regulation is vital to maintaining vascular homeostasis and healthy endothelial function, and therefore quantifying NO levels in a specific vascular area might be helpful in determining endothelial function. However, NO is a volatile substance with a...
short half-life, making its moment-by-moment quantification almost impossible (Versari et al., 2009). For this reason, NO bioavailability is usually evaluated through the measurement of its downstream effects. For example, endothelium-dependent relaxation can be measured through vascular reactivity tests (Deanfield et al., 2005), using either receptor-operated (ACh; BK; SP) stimuli in vitro and ex vivo or mechanical stimuli such as shear stress in vivo, using flow-mediated vasodilation (FMD) as the technique of assessment (John & Schmieder, 2000).

It is clear that NO and NOS play key roles in endothelial function, not only in a vasodilatory capacity, but also through a plethora of other important regulatory functions (Table 1.2). It is essential for the healthy endothelium that these two regulators are present in desirable quantities, since and imbalance in NO and NOS levels can have dire consequences for endothelial health.

1.3 Endothelial Dysfunction

When the endothelium loses its regulatory ability to maintain a balance between endothelial-derived vasodilatory and vasoconstrictory factors, it results in ED (Mudau et al., 2012). ED is characterised by a shift in the actions of the endothelium toward reduced vasodilation, a proinflammatory state, and prothrombic properties (Rajendran et al., 2013), resulting in progressive pathophysiological changes (Fig 1.5). Mechanisms by which these pathophysiological changes occur appear to be complex and multifactorial. ED is associated with most major cardiovascular risk factors (Versari et al., 2009), and is therefore a key role player in the progression to atherosclerosis and CVD. Perhaps most importantly, ED represents an initial reversible step in the development of various CVDs (Bonetti et al., 2003), making it a promising target for the prevention of such diseases when identified early (Chhabra, 2009).
Figure 1.5: The pathophysiological changes seen in a dysfunctional endothelium. In the healthy endothelium, the eNOS is responsible for most of the vascular NO production. However, eNOS becomes a potential ROS generator when in the pathological uncoupled state, due to various oxidative stresses (K. Park & Park, 2015). ACE (angiotensin-converting enzyme); Ach (acetylcholine); AT-I (angiotensin I); AT-II (angiotensin II); AT1 (angiotensin 1 receptor); BH4 (tetrahydrobiopterin); BK (bradykinin); cAMP (cyclic adenosine monophosphate); cGMP (cyclic guanosine monophosphate); ECE (endothelin converting enzyme); eNOS (endothelial nitric oxide synthase); EDHF (endothelium derived hyperpolarizing factor); ETA and ETB (endothelin A and B receptors); ET-1 (endothelin-1); L-Arg (L-arginine); L-Cit (L-citrulline); M (muscarinic receptor); O2\textsuperscript{-} (superoxide anion); ONOO\textsuperscript{-} (peroxynitrite); NADPH (nicotinamide adenine dinucleotide phosphate); NO (nitric oxide); NOX (nicotinamide adenine dinucleotide phosphate oxidase); PGH\textsubscript{2} (prostaglandin H\textsubscript{2}); PGI\textsubscript{2} (prostaglandin I\textsubscript{2}); ROS (reactive oxygen species); S1\textsubscript{B} (serotonin receptor); TP (thromboxane prostanoid receptor); TXA\textsubscript{2} (thromboxane); 5-HT (serotonin); Θ (inhibition); ⊕ (stimulation).
1.3.1 Pathophysiology

The fundamental feature of ED is the impaired or reduced NO bioavailability (Endemann & Schiffrin, 2004; Versari et al., 2009), resulting in the inability of the endothelium to initiate vasodilation in response to vasodilatory stimuli such as acetylcholine or shear stress (Mudau et al., 2012). In the presence of impaired NO bioavailability, the endothelium has to implement alternative physiological pathways in an attempt to compensate for NO deficiency (Versari et al., 2009).

ED comprises of a state of chronic “endothelial activation” (Bonetti et al., 2003), sometimes referred to as the endothelial activation–dysfunction–injury triad (Bijl, 2003). This state is characterised by a proinflammatory, proliferative, and procoagulatory milieu (Anderson, 1999), which is detrimental to the progression of CVDs, specifically all stages of atherogenesis.

Along with NO deficiency, a dysfunctional endothelium can become imbalanced in its release of other substances and mediators that are detrimental to the arterial wall, leading to excessive release of factors including endothelin-1 (ET-1), thromboxane (THA₂), prostaglandin H₂ (PGH₂), and reactive oxygen species (ROS) (Fig 1.5) (Taddei, Ghiadoni, Virdis, Versari, & Salvetti, 2003). As the endothelial function further deteriorates, the ECs exhibit reduced anti-oxidant and anti-inflammatory effects, increased vascular permeability to lipoproteins, and the increased expression of inflammatory cytokines and adhesion molecules (Libby, Ridker, & Maseri, 2002). C-reactive protein (CRP) is an example of a specific compound that moves into the tissue and causes inflammation (Devaraj, Singh, & Jialal, 2009), leading to further aggravation of the faulty endothelium.

1.3.2 Aetiology and Mechanisms of ED

Chronic exposure to cardiovascular risk factors and the harmful circulating stimuli associated with these conditions overwhelms the defence mechanisms of the vascular endothelium, hence compromising its integrity and ultimately initiating ED (Deanfield, Halcox, & Rabelink, 2007). ED has been observed to be associated with major cardiovascular risk factors, such as aging, hyperhomocysteinemia, post-menopause state, smoking, diabetes, hypercholesterolemia, hypertension, and more (Rajendran et al., 2013; Versari et al., 2009).

11
Most cardiovascular risk factors are associated with the up-regulation of intracellular oxidative stress and ROS (Cai & Harrison, 2000; K. Park & Park, 2015). Among various complex mechanisms, oxidative stress appears to be the most common underlying mechanism for the development of ED (Drexler, 1997; K. Park & Park, 2015). Oxidative stress is caused by an imbalance between the pro-oxidative and anti-oxidative molecules (Cahill & Redmond, 2016; Pennathur & Heinecke, 2007), often resulting in biomolecular damage caused by excess ROS (Halliwell, 2007). ROS is a term that includes oxygen radicals such as superoxide (O$_2^-$) and the hydroxyl radical (OH$^-$), as well as non-radical derivatives of oxygen (O$_2$) including hydrogen peroxide (H$_2$O$_2$) (Cahill & Redmond, 2016).

In most instances, the human body has an adequate supply of endogenous antioxidants, as well as antioxidants obtained from various foods to neutralise these free radicals. If the body is depleted of these antioxidants, or in the presence of multiple cardiovascular factors, injury to the endothelium and a loss in NO homeostasis can occur (Rajendran et al., 2013).

In the presence of cardiovascular risk factors, NADPH oxidases (NOX) and eNOS are often both dysregulated (Forstermann & Munzel, 2006), as a compensatory mechanism to increase NO levels and therefore all of the athero- and vaso-protective effects associated with it. The upregulation of NOX results in an increased production of ROS – specifically O$_2^-$ - in the vascular wall, with NOX being implicated as the main source for oxidative excess in the vasculature (Hamilton, Brosnan, Al-Benna, Berg, & Dominiczak, 2002). The upregulation of functional eNOS results in increased levels of NO, which is the desired effect. However, in the presence of excess O$_2^-$, NO cannot be utilised for its pro-dilatory properties, and instead O$_2^-$ combines with NO to form the highly reactive nitrogen species (RNS), peroxynitrite (ONOO$^-$) (Fig 1.6) (Forstermann & Munzel, 2006). Under physiological conditions, the enzyme sodium oxide dismutase (SOD) usually regulates O$_2^-$ levels, but in diseased states, this defensive mechanism is overwhelmed (Landmesser & Harrison, 2001). This is to be expected considering O$_2^-$ combines with NO at a rate that is three times faster than the dismutation of O$_2^-$ by SOD (Heitzer, Schlinzig, Krohn, Meinertz, & Munzel, 2001). Together, this all results in the characteristic pathophysiological reduced NO bioavailability in ED (Tomasian, Keaney Jr, & Vita, 2000). O$_2^-$, along with ONOO$^-$, also has the ability to directly inhibit the main target of NO, sGC and further enhance oxidative stress by inhibiting SOD (Munzel, Daiber, Ullrich, & Mulsch, 2005).
The consequently high levels of ONOO$^-$ are injurious to the cells, since it has the ability to oxidatively damage DNA, lipids and proteins (Mudau et al., 2012). ONOO$^-$ can undergo protonation to form peroxynitrous acid (ONOOH), or it can combine with carbon dioxide (CO$_2$) to form nitroso-peroxocarboxylate (ONOOCO$_2^-$), both of which yield tyrosine-nitrating compounds (Hurst, 2002). Via formation of these compounds, ONOO$^-$ causes the nitration of tyrosine residues of proteins, leading to formation of nitrotyrosine (Pacher, Beckman, & Liaudet, 2007), causing cell damage. In addition to being cytotoxic, ONOO$^-$ causes oxidative damage to the zinc-thiolate cluster of eNOS, resulting in the loss of the zinc ion and the formation of disulfide bonds between the enzyme monomers, and thus disruption of the binding site for BH$_4$ and L-arginine (Forstermann & Munzel, 2006). ONOO$^-$ can also oxidise tetrahydrobiopterin (BH$_4$) - which is an essential cofactor of eNOS - to the trihydrobiopterin (BH$_3$) radical. Both of the above eventually lead to eNOS uncoupling (Fig 1.8) (Forstermann & Munzel, 2006; Kuzkaya, Weissmann, Harrison, & Dikalov, 2003). Figure 1.7 gives an overview of everything discussed above.
eNOS uncoupling involves the dissociation of the ferrous-oxygen complex of eNOS. The resultant faulty eNOS enzyme can now produce $O_2^-$ from the oxygenase domain (Fig 1.8), rather than its natural product, NO (Forsterrmann & Munzel, 2006). This makes uncoupled eNOS a potential ROS genenrator (Landmesser & Harrison, 2001; K. Park & Park, 2015). In addition to ONOO$^-$-induced eNOS uncoupling, other oxidants such as $H_2O_2$ have also been shown to uncouple the enzyme (Mudau et al., 2012).
Figure 1.8: Coupled and uncoupled eNOS. (A) In the presence of sufficient levels of substrates and co-factors, and the absence of harmful reactive species, eNOS monomers will form a dimerised, coupled enzyme and produce physiological amounts of NO. (B) Decreased levels of the substrate, L-arginine and/or harmful effects exerted by increased levels of ONOO\(^{-}\), cause failure of the enzyme to dimerise, leading to the uncoupling of eNOS and the production of O\(_2^-\) instead of NO (Mudau et al., 2012).

\(\text{H}_2\text{O}_2\) is formed as a by-product of the dismutation of O\(_2^-\) by SOD (Fig 1.6). Excess production of \(\text{H}_2\text{O}_2\), along with being implicated in eNOS uncoupling, can increase eNOS expression, through transcriptional and post-transcriptional mechanisms (Forstermann & Munzel, 2006). In the presence of antioxidant enzymes such as catalase or glutathione peroxidase, \(\text{H}_2\text{O}_2\) can be dismutated into water (H\(_2\)O) and O\(_2\) (Fig 1.6). However, in the presence of transition metals – such as copper and iron - or O\(_2^-\), \(\text{H}_2\text{O}_2\) generates highly reactive hydroxyl radicals, which cause cell damage through the peroxidation of lipids (Feletou & Vanhoutte, 2006). In addition, ROS promote the contraction of VSMCs by
facilitating the mobilization of calcium and increasing the sensibility of the contractile proteins to calcium ions (Feletou & Vanhoutte, 2006). Together, all these contributors further exacerbate the overload of ROS in ED.

Inflammation is another common underlying mechanism of ED, and there seems to be a causal relationship between oxidative stress and inflammation (Karbach, Wenzel, Waisman, Munzel, & Daiber, 2014). Under physiological conditions, appropriate levels of NO, released by the healthy endothelium, control vascular inflammation. However, during ED, excess ROS aggravates vascular inflammation (K. Park & Park, 2015). Oxidative stress may amplify the vascular pro-inflammatory signaling pathways, leading to even higher levels of ROS, since inflammatory cells release $\text{O}_2^-$ (Karbach et al., 2014). ROS has been shown to upregulate adhesion and chemotactic molecules (Griendling & FitzGerald, 2003), while inflammatory markers such as C-reactive protein (CRP) have been shown to decrease eNOS activity (Venugopal, Devaraj, Yuhanna, Shaul, & Jialal, 2002). Other than oxidative stress and chronic inflammation, infection, anti-oxidant deficiency and abnormal shear stress are all mechanism implicated in ED (K. Park & Park, 2015).

1.3.3 Consequences of ED

The presence of ED can be regarded as a clinical syndrome that is associated with, and predicts an increased rate of adverse cardiovascular events (Bonetti et al., 2003). A study by Lerman & Zeiher (2005) showed an analysis of close to 2500 patients, which demonstrated that ED is strongly and independently associated with cardiovascular events (Lerman & Zeiher, 2005).

Most evidence points towards ED as one of the major pathologic changes between exposure to the cardiovascular risk factors, and the development of atherosclerotic CVD (Fig 1.9) (K. Park & Park, 2015).
Figure 1.9 shows that reduced NO bioavailability as a consequence of ED results in pro-atherosclerotic changes to the endothelial milieu (Heitzer et al., 2001). By upregulating platelet aggregation, leukocyte adhesion, and thrombosis, amongst other factors, the dysfunctional endothelium is not only an initiator, but also an important factor in the progression of atherosclerosis and CVD (Barton & Haudenschild, 2001). Thus, ED is not only a consequent feature of risk factors, but also a possible pathogenic mechanism for their onset (Versari et al., 2009).

ED is associated with most CVDs and risk factors for CVD. Amongst others, examples include hypertension, coronary artery disease, chronic heart failure, stroke, peripheral vascular disease, insulin resistance, diabetes and chronic kidney failure (Rajendran et al., 2013).

The oxidative stress associated with ED also has negative effects, independent of the ED as a whole. These include increased VSMC proliferation (resulting in thickening of the vascular wall), endothelial cell apoptosis, and increased expression and activity of matrix metalloproteinases, which are involved in the establishment of an atherosclerotic plaque (Mudau et al., 2012).
1.3.4 Prevention and Treatment of ED

The reversal of ED may be a critical point of in the prevention of atherosclerosis and CVD since ED has been implicated as the initial, reversible step in these conditions (Mudau et al., 2012). Treatment of ED involves the alteration of specific pathogenic pathways, with the goal of preserving or restoring endothelial function (K. Park & Park, 2015).

There are two classes of endothelial therapy (Fig 1.10): primary endothelial therapy involves the preservation and improvement of endothelial function in subjects without CVD, whereby lifestyle changes are made in order to reduce and prevent exposure to cardiovascular risk factors (Barton & Haudenschild, 2001). Secondary endothelial therapy involves the improvement of the dysfunctional endothelium through the treatment of underlying cardiovascular risk factors and established CVDs (Goff et al., 2014), and is applicable to both modifiable (hypertension, obesity, diabetes, dyslipidemia) and non-modifiable conditions (aging, menopause, coronary or peripheral artery disease, heart failure, end-stage renal disease) (K. Park & Park, 2015).

Figure 1.10: Therapeutic approaches to ED (K. Park & Park, 2015). CV (cardiovascular); PAOD (peripheral arterial occlusive disease).
Considering that oxidative stress is regarded by many as the main pathophysiologic mechanism leading to impaired NO bioavailability and ED (Versari et al., 2009), antioxidant substances could potentially play a substantial role in the treatment of ED. Acute studies have shown high-dose antioxidant vitamins are extremely effective in restoring normal endothelial function, however, interventional studies using oral administration of antioxidant substances including vitamin C and E have failed to provide consistent data (Virdis et al., 2004). The reason for the lack of benefit of long-term antioxidant supplementation in the setting of primary and secondary prevention, however, is not clear (Bonetti et al., 2003). Recent evidence suggests that certain antioxidants, including vitamin E, may be inappropriate to reduce oxidative stress in vivo or may even be pro-oxidant under certain conditions (Landmesser & Harrison, 2001). These mixed findings may point to the importance of testing new antioxidants or combining different antioxidant compounds in order to observe an adequate reduction of oxidative stress (Bonetti et al., 2003), a specific endpoint which will be addressed in this study (see section 1.8).

1.4 HIV/AIDS & ART

HIV is a retrovirus that was first isolated in 1983 (De Cock, 2011) and was subsequently proven to be the cause of acquired immunodeficiency syndrome (AIDS) a year later (Ruelas & Greene, 2013). Since then, the HIV/AIDS epidemic has evolved to become the greatest challenge in global health (De Cock, Jaffe, & Curran, 2012), and one of the most devastating viral pandemics in history (Freed, 2015). There are two strains of HIV, with HIV-1 being the main cause of AIDS, while HIV-2 is associated with isolated infections in central and western Africa (Sharp & Hahn, 2011). With the prolonged period between HIV infection and symptomatic AIDS – on average 11 years in adults – there was widespread transmission of HIV before recognition of the epidemic and any prevention attempts (De Cock, 2011). In the 1990s, almost 15 years after HIV/AIDS was discovered, ART was developed to combat HIV-1 infection (Laskey & Siliciano, 2014). ART has dramatically improved survival of patients with HIV/AIDS (Thienemann et al., 2013), and patients who are diagnosed and treated with ART early can experience acceptable immune recovery and expect a near-normal lifespan (Laskey & Siliciano, 2014; Thienemann et al., 2013). Although effective at suppressing viral replication, current drugs do not eradicate the virus, making treatment with ART a life-long necessity (Freed, 2015).
1.4.1 Epidemiology

As of 2016, it is estimated that there are 36.7 million people worldwide living with HIV (Fig 1.11), while around 35 million people have died from AIDS-related illnesses since the start of the epidemic (UNAIDS, 2017; WHO, 2016). Southern Africa carries the highest burden, where as few as nine countries - which account for less than 2% of the world's population - represent about one third of global HIV infections (De Cock et al., 2012).

With the rollout of ART and various prevention programmes in middle-and-low-income countries, the most affected populations are now receiving adequate treatment (Delpech, 2013). For this reason, the global prevalence of HIV has begun to plateau, and every year, there is a promising decline in newly infected individuals.

1.4.2 Virus Structure, Targets and Life-cycle

HIV-1 is a retrovirus that belongs to the lentivirus family. Infections with lentiviruses typically show a chronic course of disease, a long period of clinical latency and persistent viral replication (Rubbert, Behrens, & Ostrowski, 2007). The HIV virion has a spherical shape and a diameter of roughly 100–130 nm. The viral envelope is composed of a lipid membrane, which is derived from the host cell and contains cellular proteins, as well as about 7–12 trimeric complexes of viral envelope protein (Kirchhoff, 2013). The core antigen contains two single strands of viral RNA (Fig 1.12) with a length of close to 10,000
nucleotides. The virion contains the enzymatic equipment that is necessary for replication, namely a reverse transcriptase (RT), an integrase and a protease (Rubbert et al., 2007).

Figure 1.12: Structure of an HIV virion particle (Rubbert et al., 2007). Gp (glycoprotein); gag (group-specific antigen).

The main targets of HIV are CD4+ helper T cells, which are key regulators of the humoral and cellular immune responses (Kirchhoff, 2013). The destruction and depletion of these cells renders the body unable to defend itself against opportunistic pathogens, eventually resulting in AIDS (Laskey & Siliciano, 2014). When HIV infects an activated CD4+ T cell, it hijacks and manipulates its transcriptional and translational machinery to reproduce itself (Kirchhoff, 2013). HIV also has the ability to infect other immune cells, such as macrophages and immature dendritic cells, which may play a role in harboring the virus in a silent integrated proviral form and thus contribute to the establishment and maintenance of viral reservoirs that prevent the eradication of HIV from the human body, even during ART (Kirchhoff, 2013).

The HIV-1 life cycle is complex and can roughly be divided in an early and a late phase of replication. The early phase begins with the attachment or fusion of the virion at the cell surface, includes reverse transcription of the viral RNA to DNA amongst other steps, and ends with the integration of the proviral DNA into the host genome (Freed, 2015; Kirchhoff, 2013). The late phase of replication starts with the initiation of proviral
transcription and ends with the release of fully infectious progeny virions, which contain activated viral proteases (Kirchhoff, 2013; Laskey & Siliciano, 2014). This process is summarized below in Figure 1.13. The viral life cycle illuminates some of the challenges associated with HIV infection. For example, the HIV life cycle lasts just one to two days and the viral RT has an extremely high mutation rate of roughly 1 error per 10 000 nucleotides (Kirchhoff, 2013). Together, a short generation time, a massive virus production rate of up to 2 109 virions per day, and a large error rate (Sharp & Hahn, 2011), allow HIV to rapidly adapt to its host environment and to develop resistance against ART or immune responses (Kirchhoff, 2013).

**Figure 1.13:** A brief summary of the HIV life-cycle (Kirchhoff, 2013). Env (envelope); gp (glycoprotein); p (protein); tat (trans-activator of transcription); Rev (trans-activator involved in protein expression); gag (group-activated antigen).
1.4.4 Antiretroviral therapy (ART)

Although there is no cure for HIV, more than 30 different anti-HIV drugs have now been approved for clinical use targeting different steps in the viral life cycle (Ruelas & Greene, 2013). ART suppresses the viral load by interfering with viral replication (Laskey & Siliciano, 2014). The different classes of antiretroviral (ARV) drugs are broadly classified by the phase of the retrovirus life cycle that the drug inhibits (Fig 1.14).

![Figure 1.14: Stages of the HIV life cycle that are targeted by ARV drugs](https://scholar.sun.ac.za)

**Figure 1.14:** Stages of the HIV life cycle that are targeted by ARV drugs (Laskey & Siliciano, 2014). CCR5 (C-C chemokine receptor type 5); CXCR4 (C-X-C chemokine receptor type 4); NRTIs (nucleoside/nucleotide reverse transcriptase inhibitors); NNRTIs (non-nucleoside reverse transcriptase inhibitors); InSTIs (integrase strand transfer inhibitors); ALLINIs (allosteric integrase inhibitors); Env (envelope); CD4 (cluster of differentiation 4).

There are six main classes of ARVs (AIDSinfo, 2013; Arts & Hazuda, 2012): The first and second ARV classes are co-receptor agonists and entry inhibitors/fusion inhibitors respectively. These classes of ARVs compete with specific receptors involved in HIV attachment and therefore block the attachment and subsequent entry of the virus into the host cell. The third and fourth type, are nucleoside or nucleotide reverse transcriptase inhibitors (NRTIs/NtRTIs) and non-nucleoside reverse transcriptase inhibitors (NNRTIs). These two classes of ARVs stop the virus from converting its RNA into DNA. The fifth class of ARVs, are integrase inhibitors, which inhibit the viral enzyme integrase. Integrase is responsible for integration of viral DNA into the DNA of the infected cell. The last type of
ARVs, are Protease Inhibitors (PIs). PIs block the viral protease enzyme, which is necessary to produce mature and infectious virions.

Initially, ARV drugs were given as monotherapy, but the standard of care has since evolved to include the administration of a cocktail or combination of ARV drugs (Arts & Hazuda, 2012), in order to prevent drug resistance (Laskey & Siliciano, 2014). Combination therapy, also known as highly active antiretroviral therapy (HAART), consists of a combination of at least three ARV drugs - drawn from four of the six main ARV classes - where two or more NRTIs typically form the backbone of most regimens (Carr, 2003). In modern day treatment, co-formulations of these individual ARVs have reduced the “pill burden” and improved patient adherence (Thompson et al., 2012). There are now 19 licensed drugs in these four classes (Table 1.3), which can be combined into more than 3,000 potential HAART regimens (Carr, 2003). The firm establishment and implementation of HAART has led to major declines in opportunistic complications, making HIV infection more of a chronic disease than an acute one, and so more drugs are being used in more patients for longer periods (Carr, 2003; Ruelas & Greene, 2013).
Table 1.3: Presently available ARV drugs. Drugs in each class are listed in approximate order of approval/availability (Carr, 2003).

<table>
<thead>
<tr>
<th>Drug Class</th>
<th>Specific Drug</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nucleoside reverse transcriptase inhibitors (NRTIs)</td>
<td>Zidovudine (AZT), didanosine (ddI)<em>, zalcitabine (ddC), lamivudine (3TC)</em>, stavudine (d4T), abacavir (ABC)*, emtricitabine (FTC)</td>
</tr>
<tr>
<td>Nucleotide reverse transcriptase inhibitors (NRTIs)</td>
<td>Tenofovir disoproxil fumarate (TDF)*</td>
</tr>
<tr>
<td>Non-nucleoside reverse transcriptase inhibitors (NNRTIs)</td>
<td>Nevirapine (NVP), delavirdine (DLV), efavirenz (EFV)*</td>
</tr>
<tr>
<td>Protease inhibitors (PIs)</td>
<td>Saquinavir (SQV), indinavir (IDV), ritonavir (RTV), nelfinavir (NFV), amprenavir (APV), lopinavir/ritonavir (rLPV), atazanavir (AZV)*</td>
</tr>
<tr>
<td>Fusion inhibitors</td>
<td>Efuvirtide (ENV; T-20)</td>
</tr>
</tbody>
</table>

*Administered once daily.

1.4.5 HIV & ART in a South African Context

SA has the largest HIV-infected population globally, with an estimated 7 million people living with HIV in 2015 (Statistics South Africa, 2015). The reason for this large South African population of HIV-infected people is unclear, but various biological and social factors may play a role (De Cock, 2011). The long-term AIDS-denialism – the view that HIV is not the cause of AIDS- in SA may have also played a pivotal role, since it led to the delayed implementation of ART by the government and resulted in thousands of deaths.

SA has the largest ART programme in the world (South African National AIDS Council, 2015). In April 2014, more than 3 million people were receiving ART, which equates to 47% of people living with HIV in the country (South African National AIDS Council, 2015). This is a monumental increase compared to statistics from 12 years earlier, where only 2% of the HIV positive population in SA was receiving ART (Joint United Nations Programme
on HIV/AIDS, 2010). The extent of the HIV/AIDS epidemic and the need for increased access to ART has had a profound effect on already stretched health care systems in SA (Levitt et al., 2011). In 2013, SA changed the CD4+ cut-off value at which patients could start ART, from 200 to 350 cells/µl, making more people eligible for treatment. By the end of 2014, SA increased the level again to 500, expanding eligibility even further (South African National AIDS Council, 2015; UNAIDS, 2016). In 2015, the WHO released guidelines recommending people living with HIV be offered ART immediately following diagnosis, regardless of CD4+ count (Dutta et al., 2015). As of 2016, SA began implementing this recommendation (South African National AIDS Council, 2016). Despite all of these implementations, HIV prevalence still remains as high as 19.2% among the general South African population, although, this varies considerably between regions (UNAIDS, 2014).

1.4.5.1 SA First Line Fixed Dose Combination Therapy

In 2013, a new fixed dose combination (FDC) therapy was implemented in SA, under various trade names, including Odimune®. A single Odimune® tablet consists of two NRTIs - Emtricitabine (FTC): 200 mg / day, and Tenofovir disoproxil fumarate (TDF): 300 mg / day – as well as a NNRTI - Efavirenz (EFV): 600 mg / day (Table 1.4) (Davies, 2013). This combination constitutes the preferred first-line regimen for previously untreated patients (Meintjes et al., 2014). A once daily FDC tablet has substantially simplified the prescription, dispensing and stock management of ART drugs (Davies, 2013), as well as improved treatment adherence and the burden on patients subjected to multiple drug regimens (Meintjes et al., 2014). Table 1.4 summarizes various aspects of the different ARV drugs used in South African first line ART.
Table 1.4: Summary of ARVs used for first line FDC therapy in SA  (Arts & Hazuda, 2012; Davies, 2013; Does, Thiel, & Johnson, 2003; Meintjes et al., 2014; Palmisano & Vella, 2011; Sierra-Aragon & Walter, 2012).

<table>
<thead>
<tr>
<th>Name</th>
<th>Class</th>
<th>Recommended Dose (mg/day)</th>
<th>Mr (g/mol)</th>
<th>Mechanism of action</th>
<th>Common or severe ADR†</th>
<th>Alternative drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Emtricitabine (FTC)</td>
<td>NRTI</td>
<td>200</td>
<td>247.2</td>
<td>Cellular enzymes phosphorylate FTC to FTC-5'- triphosphate which in turn competes with the natural substrate, deoxycytidine 5'- triphosphate and is subsequently incorporated into the nascent viral DNA and leads to chain termination.</td>
<td>Palmar hyperpigmentation, hyperlactataemia/ steatohepatitis (very low potential).</td>
<td>Lamivudine (3TC).</td>
</tr>
<tr>
<td>Tenofovir disoproxil fumarate (TDF)</td>
<td>NTERT</td>
<td>300</td>
<td>287.2</td>
<td>TDF diphosphate inhibits HIV RT through competition with the natural substance deoxyadenosine 5'- triphosphate and leads to DNA chain termination after incorporation into the viral-DNA.</td>
<td>Renal failure, tubular wasting syndrome, reduced bone mineral density, hyperlactataemia/ steatohepatitis (very low potential).</td>
<td>Abacavir (ABC).</td>
</tr>
<tr>
<td>Efavirenz (EFV)</td>
<td>NNRTI</td>
<td>600 (at night) – 400 if &lt; 40kg</td>
<td>315.7</td>
<td>Diffuses into the host cell and binds next to the active site of RT which leads to a conformation change and inhibits the enzyme’s function. It is also a non-competitive inhibitor of HIV RT with respect to template, primer or nucleoside triphosphates.</td>
<td>Central nervous system symptoms (vivid dreams, problems with concentration, dizziness, confusion, mood disturbance, psychosis), rash, hepatitis, gynaecomastia.</td>
<td>Rilpivirine (RPV), nevirapine (NVP) – If EFV; RPV and NVP are contraindicated, raltegravir (RAL) or a PI can be substituted.</td>
</tr>
</tbody>
</table>

†Life-threatening reactions are included in bold. Mr (molecular weight); ADR (adverse drug reaction); mg (milligram); g (gram); mol (mole); RT (reverse transcriptase); kg (kilogram).
1.5 HIV, ART and ED/CVD

It has been established that the prevalence of CVD is augmented in HIV positive patients compared with the general population (Masiá et al., 2007), with recent studies showing the relative risk of developing CVD as high as 61% higher in HIV positive populations compared to uninfected people (Islam, Wu, Jansson, & Wilson, 2012). Factors associated with HIV infection and ART have been implicated in the premature development of atherosclerosis and coronary heart disease (CHD) (Masiá et al., 2007). CVD is the leading comorbidity and cause of death in this population (DAD Study Group, 2007; Thienemann et al., 2013), where ED is thought to be the main link between HIV infection, ART and CVD and atherosclerosis (Francisci et al., 2009), amongst other factors (Table 1.5) (de Larrañaga, Petroni, Deluchi, Alonso, & Benetucci, 2003; Wolf et al., 2002). It has been reported that enhanced oxidative stress in HIV infected patients may contribute to ED, while it has also been shown that ART and therapeutic control of HIV replication induces a pro-oxidative state (Day & Lewis, 2004; Hulgan et al., 2003).

Table 1.5: Mechanisms by which HIV and ART may adversely affect the vasculature (Dube et al., 2008).

<table>
<thead>
<tr>
<th>HIV infection</th>
<th>ART</th>
</tr>
</thead>
<tbody>
<tr>
<td>Endothelial dysfunction</td>
<td>Endothelial dysfunction</td>
</tr>
<tr>
<td>Lipid disorders associated with HIV infection</td>
<td>Increased endothelial permeability</td>
</tr>
<tr>
<td>Viral protein-related EC activation</td>
<td>Increased oxidative stress</td>
</tr>
<tr>
<td>Systemic inflammatory cytokine-chemokine dysregulation</td>
<td>Increased mononuclear cell adhesion</td>
</tr>
<tr>
<td>Direct HIV infection of endothelium and VSMCs</td>
<td>Insulin resistance</td>
</tr>
<tr>
<td>Enhanced atheroma formation by activated macrophages</td>
<td>Accelerated lipid accumulation in vessel wall</td>
</tr>
<tr>
<td>Prothrombotic state</td>
<td>Persistent inflammation and immune activation</td>
</tr>
<tr>
<td></td>
<td>Impaired response to vascular injury</td>
</tr>
</tbody>
</table>
However, it is still debated whether cardiovascular complications are a consequence of the HIV infection itself or due to the long-term use of ART (Francisci et al., 2009). Certain studies have shown that it may be the ART that is responsible for the increased risk for CVD (Obel et al., 2007), with one study in particular showing that the risk of CVD for people living with HIV receiving ART was found to be two times greater than the risk for HIV positive patients who were treatment-naïve (Islam et al., 2012). Other studies contest this evidence, suggesting HIV infection itself and not ART is the pathological initiator (Baker et al., 2011; Francisci et al., 2009). The Strategies for Management of Anti-Retroviral Therapy (SMART) study, showed that persons undergoing episodic ART had an increased risk of cardiovascular events identical to that of patients undergoing continuous therapy (Strategies for Management of Antiretroviral Therapy (SMART) Study Group, 2006). Some studies have even shown that ART actually has a protective role in terms of risk for ED and CVD (Arildsen, Sørensen, Ingerslev, Østergaard, & Laursen, 2013; Francisci et al., 2009; Grubb et al., 2006; Solages et al., 2006; Torriani et al., 2008; Wolf et al., 2002), suggesting rather that viral load, CD4+ cell count and cardiovascular risk factors are responsible for the endothelial function impairment observed in other studies (van Wijk et al., 2006; Wolf et al., 2002). It is important to note however, that most studies showing a cardioprotective role for ART, were performed in treatment-naïve patients, initiating short term ART treatment (up to 24 weeks) (Torriani et al., 2008). Therefore it is uncertain if the improvements were due to ART, suppression of viremia, or changes in immune activation (Torriani et al., 2008). Unfortunately, it is still unclear how to distinguish the effects of a chronic HIV infection from those of ART (Skowyra, Zdziechowicz, Mikula, & Wiercińska-Drapało, 2012), making a consensus more difficult to obtain.

1.5.1 ART induced ED/CVD

Although ART has dramatically reduced HIV-associated morbidity and mortality (Jiang et al., 2007), life-long treatment may expose patients to an increased risk for developing CVD (DAD Study Group, 2007; SMART study group, 2006). With recent consensus guidelines recommending the initiation of ART as early as possible in the course of HIV infection, the average duration of exposure to ART is increasing, and therefore complications associated with treatment are also expected to become more common (Choi et al., 2011). Despite all the confounding reports, the vast majority of trials indicate that ART can lead to ED (Skowyra et al., 2012). Cardiovascular complications are considered a new challenge for HIV patients receiving ART with numerous studies investigating the effects of ART on the
vasculature indicating that ART may impair the function of the vascular endothelium (Blanco et al., 2006; Shankar, Dubé, Gorski, Klaunig, & Steinberg, 2005; Stein et al., 2001). The Data Collection on Adverse Events of Anti-HIV Drugs (DAD) study group reported that ART was associated with a 26% increase in the rate of myocardial infarction (MI) per year of exposure to ARV drugs (DAD Study Group, 2007). This finding lead to the theory that duration of exposure to ART may be one of the main factors associated with risk of developing CVD, which has been confirmed in other studies (Islam et al., 2012).

The major underlying mechanism is thought to be ART-induced NO serum level reduction, as well as increased ROS and cholesterol (Skowyra et al., 2012). ART-induced ROS production is thought to be of mitochondrial origin, likely resulting from mitochondrial dysfunction and leakage of electrons from a dysfunctional mitochondrial electron transport chain (Jiang et al., 2007). There is controversy however; surrounding which ART class has the highest degree of associated risk of developing CVD.

1.5.1.1 Protease Inhibitors

The majority of studies investigating ART have demonstrated that PI's are associated with increased cardiovascular risk. PIs are thought to directly induce ED by damaging mitochondrial DNA of ECs (Skowyra et al., 2012). An early study documented severe ED in patients who received long-term, PI–based ART (mean duration of ART was 70 months, including 31 months of therapy with a PI) but not in those receiving ART without a PI (Stein et al., 2001). In contrast, another early observational study of hospitalisation rates in Northern California, Klein et al. (2002) found that PIs did not tend to increase the rates of hospitalisations for CHD among persons living with HIV. Increased risk of MI in HIV patients receiving PI-based ART has been reported by multiple cohorts (DAD Study Group, 2007; Holmberg et al., 2002; Lang et al., 2010). Yeboah et al. (2007) demonstrated that the administration of a PI impaired endothelial function in healthy men, while several other studies have identified cellular or molecular mechanisms of ED associated with the use of PIs (Grubb et al., 2006; Wang, Chai, Yao, & Chen, 2007; Zhong et al., 2002). Clinical and experimental models have shown that PI-induced ED appears to be mediated by reduced NO production (Shankar et al., 2005), due to reduced expression of eNOS (Fu, Chai, Yao, & Chen, 2005) and increased ROS (Dube et al., 2008).
1.5.1.3 Nucleoside Reverse Transcriptase Inhibitors

More recent studies have also implicated NRTI-based ART in playing a damaging role (Carr, 2003; Choi et al., 2011; Gallant, Parish, Keruly, & Moore, 2005; Jiang et al., 2007; Sutliff et al., 2002; Young et al., 2007), with far less literature being available regarding the specific effects of NNRTI-based ART (Hulgan et al., 2003; Stein et al., 2001). This is of particular relevance for South African first line ART, which is composed of two Marti’s and one NNRTI. NRTI-based ART use is also associated with an increased risk of CVD, but not to the same extent as PI-based ART (Islam et al., 2012). Mitochondrial toxicity is recognised as the most common side effect of NRTIs (Sutliff et al., 2002), and is associated with the development of a number of tissue-specific complications (Lewis & Dalakas, 1995). Mitochondrial dysfunction is thought to be the result of two interrelated pathophysiological mechanisms that result from defective oxidative phosphorylation, namely increased production of mitochondrial ROS and reduced energy production (Lewis, Copeland, & Day, 2001). Furthermore, the production of free radicals has been implicated in mitochondrial DNA damage and dysfunction in ECs (Ballinger et al., 2000). A study by Sutliff et al. (2002) demonstrated that NRTIs profoundly impaired endothelium-dependent relaxation in murine aortas, with increased O$_2^-$ levels being implicated as the mediator of these adverse effects. This finding has been supported by numerous other studies that have found a link between NRTI toxicity and increased free radical levels (de la Asuncion et al., 1998; Jiang et al., 2007; Jiang, Hebert, Zavecz, & Dugas, 2006). Together, these mechanisms are thought to be responsible for the association of NRTIs with accelerated vascular disease such as platelet activation, hyperlipidemia, T-lymphocyte activation, abnormal endothelial function and vascular inflammation (Choi et al., 2011). In contrast, a recent meta-analysis of clinical trial studies found that NRTIs were not associated with a greater risk of MI or major CVD events, despite reports from other cohort studies (Cruciani et al., 2011).

It is important to note that majority of the studies reported in the literature are conducted using one or two specific PIs or NRTIs. Therefore one should be wary when making conclusions, since adverse effects discovered in a particular drug should not generally be considered representative of the entire drug class (Dube et al., 2008). It is thus very important to expand the literature and report findings on all ARV drugs currently in use, and in all the different classes of ART.
1.6 Melatonin

Melatonin is a hormone found in all vertebrates, and almost ubiquitously present in bacteria, protozoa, plants, fungi and invertebrates (Hardeland & Poeggeler, 2003). It is rhythmically secreted by the pineal gland, and involved in regulation of circadian and, sometimes, seasonal rhythms (Reiter, 1993). Melatonin is an indoleamine, which contributes to its amphiphilicity, which means it has the ability to enter any cell, compartment or body fluid (Hardeland, Pandi-Perumal, & Cardinali, 2006). During melatonin synthesis, its precursor tryptophan, is taken up from the blood and converted, via 5-hydroxytryptophan, to serotonin. Serotonin is then acetylated to form N-acetylserotonin by arylalkylamine N-acetyltransferase (AA-NAT), which, in most cases, represents the rate-limiting enzyme. N-acetylserotonin is then converted into melatonin by hydroxyindole O-methyltransferase (Fig 1.15) (Pandi-Perumal et al., 2006; Sprenger, Hardeland, Fuhrberg, & Han, 1999).

![Pathways of indolic catabolism involved in the formation of melatonin](Pandi-Perumal et al., 2006)

**Figure 1.15:** Pathways of indolic catabolism involved in the formation of melatonin (Pandi-Perumal et al., 2006).
Pineal melatonin production exhibits a circadian rhythm, with a low level during daytime and high levels during night (Pandi-Perumal et al., 2006). Melatonin is able to retrain the circadian rhythm through its short biological half-life, whereby serum levels of approximately 0.4 nM peak at night (Brzezinski, 1997). This circadian rhythm persists in most vertebrates, irrespective of whether the organisms are active during the day or during the night (Borjigin, Li, & Snyder, 1999; Claustrat, Brun, & Chazot, 2005). Other than regulation of circadian rhythms, melatonin also appears to play a role in immunomodulation, reproduction, tumor growth, and aging (Brzezinski, 1997). Most importantly, melatonin has also been shown to exhibit strong antioxidative properties (Hardeland et al., 2006).

1.6.1 Antioxidant Actions

Melatonin has been reported to have both direct and indirect antioxidant properties (Fig 1.16) (Bonnefont-Rousselot & Collin, 2010; Hardeland, 2005; Tan et al., 2002). Direct antioxidant properties of melatonin include the scavenging of ROS (Poeggeler et al., 2002), while indirectly, melatonin upregulates antioxidant enzymes (Hardeland, 2005). The antioxidant activity of melatonin has been reported at both physiological and pharmacological concentrations (Galano, Tan, & Reiter, 2011; Reiter, Tan, & Maldonado, 2005), although pharmacological doses of exogenously administered melatonin typically provide greater protection from large quantities of free radicals (Tan, Manchester, Terron, Flores, & Reiter, 2007). As a whole, the antioxidant properties of melatonin suggest that it could have significant potential in treating chronic diseases associated with oxidative stress (Fig 1.16).
Figure 1.16: Direct and indirect actions of melatonin in an antioxidative capacity (Bonnefont-Rousselot & Collin, 2010). Directly, melatonin acts as an antioxidant, where it has the ability to directly neutralise reactive species. Indirectly, melatonin up-regulates other antioxidant enzymes that in turn, are capable of removing various reactive species. GSH-Px (glutathione peroxidase); ROS (reactive oxygen species); SOD (superoxide dismutase).
1.6.1.1 Free radical Scavenging

There is a vast amount of literature documenting melatonin’s ability to directly scavenge both ROS and RNS (Allegra et al., 2003; Poeggeler et al., 2002; Reiter, Tan, Manchester, & Qi, 2001; Tan et al., 2002). Specific species removed by melatonin include OH–, \( \text{H}_2\text{O}_2 \), singlet oxygen, \( \text{O}_2^- \), NO and ONOO–.

Melatonin is able to neutralise OH– by trapping OH– in one of its indole rings (Tan et al., 2002). \( \text{H}_2\text{O}_2 \) and singlet oxygen - an energy-rich form of \( \text{O}_2 \) - are thought to be removed through their reaction with melatonin is the presence of the substrate \( \text{N}1\text{-acetyl-N}2\text{-formyl-5-methoxykynuramine (AFMK)} \) (Tan et al., 2000), although this has only been demonstrated in a pure chemical system and whether intracellular melatonin neutralizes \( \text{H}_2\text{O}_2 \) in this manner is unknown (Reiter et al., 2003). Surrounded by more uncertainty, is the efficacy of melatonin in neutralizing the \( \text{O}_2^- \), which is poorly defined, and still particularly unclear in vivo. Far more is known regarding melatonin’s interaction with NO. Blanchard et al. (2000) initially showed that melatonin is only capable of interacting with NO in the of presence of \( \text{O}_2 \), suggesting that melatonin may rather react with a molecule derived from NO, possibly ONOO– (Reiter et al., 2003). See Table 1.6 for a summary.

1.6.1.2 Antioxidant Upregulation

Antioxidant enzymes represent a first line of defence against toxic oxidant reactants by metabolizing them to innocuous by-products (Rodriguez et al., 2004). The main antioxidant enzymes upregulated by melatonin are superoxide dismutase (SOD), catalase (CAT), glutathione peroxidase (GPx), glutathione reductase (GRd) and tripeptide glutathione (GSH) (Bonnefont-Rousselot & Collin, 2010; Hardeland, 2005; Hung et al., 2013; Tengattini et al., 2008).

As previously mentioned on page 13, SOD converts \( \text{O}_2^- \) to \( \text{H}_2\text{O}_2 \), thereby decreasing the formation of ONOO– (Mudau et al., 2012). SOD works in conjunction with other antioxidant enzymes such as CAT and GPx, which are also capable of removing \( \text{H}_2\text{O}_2 \). (Tengattini et al., 2008). CAT promotes the conversion of \( \text{H}_2\text{O}_2 \) to \( \text{H}_2\text{O} \) and \( \text{O}_2 \), while GPx converts peroxides to \( \text{H}_2\text{O} \) using GSH as a substrate (Valko, Rhodes, Moncol, Izakovic, & Mazur, 2006). Melatonin further facilitates this reaction by maintaining high intracellular levels of GSH. Besides upregulating antioxidants, melatonin can also downregulate pro-oxidative enzymes like NOS, which in turn decreases NO production, and decreases the amount of NO available to react with \( \text{O}_2^- \), ultimately decreasing ONOO– levels (Aydogan, Yerer, &
Goktas, 2006; Karbownik & Reiter, 2000). All these effects of melatonin are summarised in Table 1.6 and Figure 1.17 below.

This regulation of antioxidants and oxidants by melatonin is controlled through messenger-ribonucleic acid (mRNA) (Hung et al., 2013). When melatonin binds to its receptor, it stimulates the phospholipase C pathway. The consequent increase in calcium concentration leads to the phosphorylation of protein-kinase C (PKC). PKC activates protein/activation transcription factor cAMP responsive element binding protein and activating transcription factor (CREB-ATF) (Tengattini et al., 2008). This pathway modulates immediate early gene transcription and consequently gene transcription regulation and antioxidant enzyme levels.

The effects of melatonin on the mitochondria has become another recent research focus. It is well established that the opening of the mitochondrial permeability transition pore (mtPTP) causes mitochondrial swelling and rupture of the outer mitochondrial membrane, leading to the release of pro-apoptotic factors such as cytochrome c (Petrosillo et al., 2009). Melatonin appears to have the ability to safeguard the respiratory electron flux, reduce oxidant formation by lowering electron leakage, and inhibit the opening of the mtPTP (Hardeland, 2005). A study by Petrosillo et al. (2009) showed that melatonin treatment had the ability to completely inhibit mitochondrial cytochrome c release in the hearts of male Wistar rats during ischemia-reperfusion. The same study also found that melatonin treated hearts were also less sensitive to mtPTP opening, demonstrating overall the protective effect of melatonin through inhibiting the opening of the mtPTP.
Table 1.6: Effects of melatonin on ROS/RNS and oxidant enzymes (Tengattini et al., 2008). ROS (reactive oxygen species); RNS (reactive nitrogen species); \( \downarrow \) (decrease); \( \uparrow \) (increase).

<table>
<thead>
<tr>
<th>Action</th>
<th>Target</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scavenging</td>
<td>ROS</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Hydrogen peroxide</td>
<td>( \downarrow )</td>
</tr>
<tr>
<td></td>
<td>Hydroxyl radical</td>
<td>( \downarrow )</td>
</tr>
<tr>
<td></td>
<td>Singlet oxygen</td>
<td>( \downarrow )</td>
</tr>
<tr>
<td>RNS</td>
<td>Nitric oxide</td>
<td>( \downarrow )</td>
</tr>
<tr>
<td></td>
<td>Peroxynitrite anion</td>
<td>( \downarrow )</td>
</tr>
<tr>
<td>Up-regulation</td>
<td>Antioxidant enzymes</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Superoxide dismutase</td>
<td>( \uparrow )</td>
</tr>
<tr>
<td></td>
<td>Catalase</td>
<td>( \uparrow )</td>
</tr>
<tr>
<td></td>
<td>Glutathione peroxidase</td>
<td>( \uparrow )</td>
</tr>
<tr>
<td></td>
<td>Glutathione reductase</td>
<td>( \uparrow )</td>
</tr>
<tr>
<td></td>
<td>Tripeptide glutathione</td>
<td>( \uparrow )</td>
</tr>
<tr>
<td>Down-regulation</td>
<td>Pro-oxidative enzymes</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nitric oxide synthase</td>
<td>( \downarrow )</td>
</tr>
</tbody>
</table>
1.6.2 Melatonin in ART induced ED

As discussed earlier, it has been established that the main cause of ART induced ED is oxidative stress due to excessive generation of ROS and RNS, and an imbalance in anti- and pro-oxidative substances. For these reasons, it has been suggested that the treatment or prevention of ART induced ED might be achieved through the use of antioxidants (Dube et al., 2008). A study by De la Asuncion et al. (1998) showed that treatment with antioxidant vitamins significantly reduced the development of NRTI-mediated cardiac myopathies, while on the other hand, other studies have shown antioxidant vitamins to have no effect (Sutliff et al., 2002), while the clinical use of antioxidants to ameliorate ED in patients receiving ART is not well documented (Dube et al., 2008). Therefore, more research is needed in order to determine whether antioxidants other than vitamins can be effective in combating ART induced ED.

Melatonin is the optimal antioxidant since it has both the capacity to scavenge radicals as well as upregulate other antioxidants and down-regulate pro-oxidative enzymes (Bonnefont-Rousselot & Collin, 2010; Hardeland, 2005; Tan et al., 2002). Melatonin is a
particularly favourable putative therapeutic agent for the treatment of ART induced ED since it has the ability to cross physiological barriers and enter cells, unlike other antioxidants whose actions are limited because of their solubility that limits their partitioning between intra- and extra-cellular compartments (Bonnefont-Rousselot, Collin, Jore, & Gardès-Albert, 2011). Melatonin also offers protection against radical-induced mitochondrial damage, which is an important attribute considering mitochondria have been implicated as one of the main sources of ART-induced ROS generation (Jiang et al., 2007; Lewis et al., 2001). Melatonin has the ability to inhibit the mitochondrial production of superoxide anion and hydrogen peroxide that directly activates the mtPTP - a critical event in mitochondrial-mediated cell death (Bonnefont-Rousselot et al., 2011). The inhibition of the mitochondrial production of \( \text{O}_2^- \) and \( \text{H}_2\text{O}_2 \) by melatonin has been shown to maintain the efficiency of oxidative phosphorylation and ATP synthesis while increasing the activity of the respiratory complexes (López et al., 2009). Therefore, in theory, melatonin has promising potential as a therapeutic agent, which can be co-administered with ART, however, further research is needed.

1.7 Conclusion

The wide spread use of ART has dramatically reduced HIV/AIDS related morbidity and mortality. With the early commencement of ART, patients are spending longer periods of their lives on these drugs, and due to ART-associated increased life expectancy of these patients, there is a large population of people currently living with HIV, receiving ART. It has been established that long-term or life-long exposure to ART has been associated with increased risk of developing ED. ED is a critical factor in the initiation and of progression CVD, but with early detection and treatment, ED can be reversed. The underlying mechanisms of ART induced ED are still unclear, but the main cause seems to be increased levels of ROS/RNS resulting in oxidative stress and damage. It has been proposed that antioxidant treatment could be a potential solution; with melatonin possessing all the desired properties needed to reduce and potentially reverse ART induced ED. However, literature documenting the specific relationship between melatonin and ART treatment is scarce, and therefore further research is required.
1.8 Problem Identification and Study Aims

1.8.1 Problem Identification

There is a substantial amount of literature documenting the relationship between HIV, ART and ED (Day & Lewis, 2004; Francisci et al., 2009; Masià et al. 2007). These studies are limited by the inherent difficulty in differentiating between direct drug-based effects and potential HIV effects, secondary host immunological responses, or metabolism-mediated effects on the endothelium (Grubb et al., 2006). There is a lack of studies that investigate the effects of ART alone, with less than a hand-full of studies to date looking at the effects of South African first line FDC ART (EFV + FTC + TDF) on endothelial health (Charania, 2017; Imperial, 2017; Mashele et al., 2016). With the South African health authorities increasing the output and accessibility of ART, and the documented unfavorable effects of NRTI's in particular on the endothelium and cardiovascular system (Choi et al., 2011; Jiamg et al., 2007; sutliff et al., 2002), it is important to assess this relationship. Since ART is a life-long therapy, it is important to find a potential co-treatment that will not interact with the effects of the ARV drugs themselves, as well as a therapy which can be used long term without any negative side effects. Melatonin meets these criteria, since it has the ability to target all the suggested mechanisms of ROS and RNS production caused by ART. Furthermore, melatonin is a naturally occurring hormone that has shown no toxicity, even at supra-pharmacological levels (Guardiola-Lemaitre, 1997; Tan et al., 2007). All these advantages of melatonin supplementation make the antioxidant hormone a preferential choice over other antioxidative supplements. Therefore, we propose that melatonin may have the potential to ameliorate potential endothelial damage caused by the first line FDC ART drug currently used in SA.

1.8.2 Main Study Aim

To investigate the effects of melatonin supplementation during first line FDC ART: an in vitro, ex vivo and in vivo approach.

1.8.2.1 Specific in vitro Study Objectives

- Determine the individual optimum concentrations of both melatonin and first line FDC ART treated aortic endothelial cells (AECs) at which the greatest differences are seen when compared to untreated AECs, with respect to:
  - NO production
  - RNS production
Cell viability

Use the optimum concentrations obtained in the dose-response experiments above, to treat AECs with melatonin and ART in combination, in order to analyse any potential synergistic/antagonistic interactions between the two drugs with respect to:

- NO production
- RNS production
- Cell viability

- Examine various protein activation and/or expression levels of important vascular signalling molecules in Melatonin- and/or ART-treated AECs.
- Examine antioxidant capacity exhibited by Melatonin- and/or ART-treated AECs.

1.8.2.2 Specific ex vivo and in vivo Study Objectives

- Determine the acute effects of melatonin and ART administration on the vascular reactivity of untreated, control male Wistar rat aortas.
- Determine the chronic effects of melatonin and ART administration on the vascular reactivity of male Wistar rats that have been treated for 8 weeks with melatonin and/or ART.
- Examine various protein activation and/or expression levels of important vascular signalling molecules in the aortic tissue of male Wistar rats that were treated for 8 weeks with melatonin and/or ART.
- Examine antioxidant capacity exhibited in the aortic tissue of the rats subjected to the same treatment as above.
2 - Materials and Methods: *In vitro* studies

Chapter 2 contains the materials and methods used for the *in vitro* aortic endothelial cell (AEC) culture experiments. The results for these studies can be found in Chapter 4.

### 2.1 Aortic Endothelial Cell Culture

For all cell culture experiments, a sample size of 1 was considered as an experiment conducted on a single day with a single batch of cells (biological replicate). The cells of the single experiment \((n = 1)\) were divided amongst different plates and randomly subdivided into the various treatment groups \((2 - 3 \text{ plates per treatment group} = \text{technical replicates})\). The mean value of the technical replicates constitutes the value for a single sample \((n = 1)\). This process was then repeated on a different day with a new batch of cells in order to increase the sample size. A sample size of \(3 - 4\) biological replicated \((n = 3 - 4)\) was considered sufficient, since it is the standard sample size optimised and used by our laboratory (Genis, 2014; Mudau; 2012; Strijdom et al., 2008), as well as used in other publications using aortic endothelial cells (Fulton et al., 2002; Mondal et al., 2013).

#### 2.1.1 Materials

- AECs (adult rat): Purchased from *VEC Technologies* (University at Albany Foundation, 1 University Pl, Rensselaer, NY 12144, USA).
- Attachment factor and trypsin (500 BAEE units’ trypsin / 180 μg EDTA•4Naper ml in Dulbecco’s PBS): *Life Technologies* (Carlsbad, California, USA).
- Foetal bovine serum (FBS): *Biochom-Biotec*.
- Dimethyl sulfoxide (DMSO): *Sigma-Aldrich* (St Louis, Mo, USA).

#### 2.1.2 Passaging

Pure primary adult rat AECs were received from the USA in culture in 75 cm\(^3\) flasks and further grown in EGM-2 growth medium in our own laboratory until fully confluent. EGM-2 growth medium was supplemented with 10% FBS; heparin; long chain human insulin-like growth factor (R\(^3\)-IGF-1); recombinant human epidermal growth factor (rhEGF); antibiotic GA 1000 (contains gentamicin and amphotericin); vascular endothelial growth factor (VEGF); ascorbic acid; recombinant human fibroblastic growth factor B (rhFGF-B) and...
hydrocortisone in accordance with the manufacturer’s instructions. The cells were grown in a standard tissue culture incubator (Forma Series II, Thermo Electron Corporation, Waltham, MA, USA) at 37 °C in a 40 – 60% humidified, 5% CO₂ atmosphere, and received fresh medium every second day. The primary AECs were grown to various “Parent (P)-generations” (see Figure 2.1) and then suspended in a freezing medium (90 % FBS, 5 % growth medium and 5 % DMSO) and stored in liquid nitrogen for future use. Cultures between the 4th and 7th generation were used for experiments. These parent cells were removed from liquid nitrogen, allowed to thaw and given fresh medium. Once the new cultures reached confluency, the cells were washed with phosphate-buffered saline (PBS) and incubated with pre-warmed (37 °C) trypsin until cells detached from the bottom of gelatin-containing attachment factor-coated 35 mm petri dishes (approximately 3 minutes). Using a 1 ml pipette, the detached cells were then rapidly transferred to a 15 ml conical tube containing fresh growth media, which deactivates the trypsin and prevents the cells from lysing. The tube containing the cells was then centrifuged for 3 minutes at 1000 revolutions per minute (rpm). The supernatant was aspirated and pellet re-suspended in fresh growth medium. The media containing the cells was then split between new 35 mm petri dishes that had been pre-incubated with attachment factor for at least 1 hour prior to the passage. Passaging to the next generation was performed at a 1:2 ratio. A summary of the passaging process can be found in Figure 2.1.
Figure 2.1: Passaging and cell aliquot storage procedures (Genis, 2014).
2.2 Experimental Groups and Protocols

All experiments were performed on confluent cells. The close cell-to-cell contact minimises possible cell cycle variability when evaluating experimental results. The cells remain morphologically and functionally viable, however since they enter cell-cycle arrest, they are no longer proliferative. For all cultures, petri dishes were randomly assigned to respective control and experimental groups. Cells were serum starved before being treated in order to induce cell cycle synchronization, which enhances the cells’ responses to experimental treatment (Gerber et al., 1998; Russell & Hamilton, 2014), in otherwise healthy cells. For cells to be serum starved, they were incubated in medium containing 0 % FBS for 24 hours prior to treatment with the various drugs.

2.2.1 Melatonin and Vehicle

For melatonin dose-response experiments, 23.2 mg of melatonin (Sigma Aldrich, St Louis, MO, USA) was dissolved in 150 µl methanol (MeOH) vehicle. This solution was further dissolved in a total volume of 10 ml by adding 9.850 ml of distilled water (dH₂O), thus creating a 10 mM melatonin stock. The 10 mM melatonin stock was further diluted in dH₂O to make various concentrations. Freshly made melatonin stock was used for every experiment, just before treatment. Since melatonin is highly light sensitive, the lights were always turned off during preparation, dilution and treatment with melatonin. All tubes containing melatonin were also covered in tin foil for extra protection from the light. Vehicle preparations followed the same protocol as melatonin preparation, with the omission of melatonin.
Melatonin dose-response experiments consisted of the following groups:

- **Absolute Control (AC)**
  - The AC group is a control group that does not receive any fluorescent probe and therefore represents the auto-fluorescence of cells.

- **Control (C)**
  - The C group in an untreated group where normal media is added to the plate, and the plate is kept in the same conditions as the treated plated over the treatment period.

- **Positive Control (+C)**
  - The +C group is treated with a product that is known to induce the expected effect and therefore validates the experimental procedure.

  - 1 nM Melatonin (1 nM Mel)
  - 1 nM Vehicle (1 nM Veh)
  - 1 µM Melatonin (1 µM Mel)
  - 1 µM Vehicle (1 µM Veh)
  - 10 µM Melatonin (10 µM Mel)
  - 10 µM Vehicle (10 µM Veh)

Refer to Appendix A for detailed melatonin and vehicle concentration calculations.

### 2.2.2 ART and Vehicle

South African first line FDC ART consists of three individual ARV drugs, namely efavirenz (EFV), emtricitabine (FTC), and tenofovir (TDF). For experimental purposes, drugs were purchased in their individual ARV form from SantaCruz Biotechnology (WhiteHead Scientific in SA). The drugs were received in 10 mg powder form and dissolved in various solvents/vehicles.

For ART dose-response experiments, three concentrations were chosen for each individual ARV component (EFV: 5 µM; 8 µM; 12 µM; FTC: 5 µM; 7.5 µM; 10 µM; TDF: 80 nM; 400 nM; 1 µM). The various concentrations were chosen based on literature documenting the use of these specific ARV drugs in cells and rats (Bertrand & Toborek, 2015; Borroto-Esoda, Vela, Myrick, Ray, & Miller, 2006; Bousquet, Pruvost, Guyot, Farinotti, & Mabondzo, 2009; De Pablo et al., 2012; Glover et al., 2014; Jamaluddin, Lin, Yao, & Chen, 2010). The average serum concentration found in humans after taking the recommended dose of the relevant individual ARV drug was also taken into consideration when deciding on the final concentrations (Kearney, Flaherty, & Shah, 2014).
2004; Luber et al., 2010; Molina et al., 2004; Villani et al., 1999). The three concentrations were either classed as a low, mid or high concentration for the purposes of this study. For experimental treatment, the individual ARV drug components (EFV; FTC and TDF) were combined to form a drug cocktail similar to the first line FDC ART drug regimen. The three drugs were combined in such a way that the low concentration of each individual ARVs were combined to form one low concentration FDC ART regimen. The same was applied for the mid and high concentrations (Fig 2.2). See below for the specific concentrations used in the low, mid and high concentration groups.

ART dose-response experiments consisted of the following groups:

- Absolute Control (AC)
  - The AC group is a control group that does not receive any fluorescent probe and therefore represents the auto-fluorescence of cells.
- Control (C)
  - The C group is an untreated group where normal media is added to the plate, and the plate is kept in the same conditions as the treated plated over the treatment period.
- Positive Control (+C)
  - The +C group is treated with a product that is known to induce the expected effect and therefore validates the experimental procedure.
- Low concentration ART (Low ART)
- Low concentration Vehicle (Low Veh)
- Mid concentration ART (Mid ART)
- Mid concentration Vehicle (Mid Veh)
- High concentration ART (High ART)
- High concentration Vehicle (High Veh)

Refer to Appendix B for detailed ARV and vehicle concentration calculations.
2.2.2.1 EFV

The 10 mg EFV powder was dissolved in 1.5 ml of MeOH vehicle to create an EFV stock solution (21.119 mM). This stock was then divided into 40 µl aliquots and frozen away at -20 °C until needed. For experiments, the required amount of aliquot was thawed and added in specific quantities to growth medium to yield the desired final concentration of ART. Vehicle preparations followed the same protocol, with the omission of the actual drug.

2.2.2.2 FTC

The 10 mg FTC powder was dissolved in 4 ml of dH₂O vehicle to create an FTC stock solution (10.111 mM). This stock was then divided into 65 µl aliquots and frozen away at -20 °C until needed. For experiments, the required amount of aliquot was thawed and added in specific quantities to growth medium to yield the desired final concentration of ART. Vehicle preparations followed the same protocol, with the omission of the actual drug.

2.2.2.3 TDF

The 10 mg TDF powder was dissolved in 40 ml of dH₂O vehicle to create TDF stock solution A (0.3934 mM). TDF stock solution A was then divided into 350 µl aliquots and frozen away at -20 °C until needed. For experiments, the required amount of aliquot was thawed, after which a stock solution B (50 µM) was made, in order to obtain the lowest concentration that TDF was used in. 127.1 µl TDF of stock solution A was added to 872.9 µl of dH₂O to create TDF stock solution B. Both stocks were added in specific quantities, where applicable, to growth medium to yield the desired final concentration of ART. Vehicle preparations followed the same protocol, with the omission of the actual drug.

Figure 2.2: Individual ARV drug concentrations that were combined to make-up the three fixed dose combination ART treatment groups.
2.2.3 Combined drugs and Vehicles

For the combination studies, one concentration of melatonin and one concentration level for ART was selected, based on dose-response results (See Chapter 4, sections 4.1.1 and 4.1.2). We chose to use 1 nM melatonin along with the high concentration ART, based on the fact that these concentrations resulted in the greatest change in the investigated variable (Fig 4.6; 4.13; 4.14 & 4.15). The same stock and vehicle preparation methods were used in the combination studies as was described above, to achieve the desired concentrations, with the combined vehicle containing a mixture of MeOh (for melatonin and EFV) and dH2O (for FTC and TDF).

2.3 Plate Reader Analyses

2.3.1 Materials

- 4,5-diaminofluorescein-2/diacetate (DAF-2/DA): Calbiochem (San Diego, CA, USA).
- Dihydrorhodamine-1, 2, 3 (DHR-123): Sigma-Aldrich (St Louis, Mo, USA).
- Diethylamine NONOate diethylammonium salt (DEA/NO): Sigma-Aldrich (St Louis, Mo, USA).
- Propidium iodide (PI) solution: Biochom-Biotech (San Diego, CA, USA).
- Authentic peroxynitrite: Millipore (Billerica, MA, USA)
- All other chemicals and buffer reagents: Merck (Damstadt, Germany).

2.3.2 Methods

At the final passage, AECs were re-seeded onto Vision Plates™ (clear bottom, black-walled 24-well plates) (4titude, Surrey, UK) and grown to confluency for experimentation. Upon the establishment of confluence, the cells were serum starved for 24 hours (see section 2.2) and treated with the various drugs. After 1- or 24-hours of drug treatment, the cells were rinsed with pre-warmed (37 °C) PBS and treated with the applicable fluorescent probe (in the dark). The fluorescence intensity was measured using a FLUOstar Optima microplate reader (BMG Labtech, Germany). Figure 2.3 A gives an example of an experimental layout of a single sample (n = 1) on a given day, while the general protocol followed for each experiment is summarised in Figure 2.3 B. Table 2.1 shows the specific excitation and emission wavelengths used on the microplate reader for the various
probes. Data generated by the microplate reader was stored and analysed with MARS Omega Data Analysis (software version 1.30; BMG Labtech, Germany).

**Table 2.1:** Excitation and emission wavelengths used for the various probes on the microplate reader

<table>
<thead>
<tr>
<th>Settings</th>
<th>DAF</th>
<th>DHR</th>
<th>PI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Excitation</td>
<td>485</td>
<td>485</td>
<td>544</td>
</tr>
<tr>
<td>Emission</td>
<td>520</td>
<td>520</td>
<td>640</td>
</tr>
</tbody>
</table>

Cells are passaged to the appropriate

On the final passage, the cells are passaged

On establishment of confluence, the individual wells are split into the various treatment groups and treated. See below for an example of the layout used for treatment groups on a 24-well plate:

<table>
<thead>
<tr>
<th>TG 1 (Tech. rep. #1)</th>
<th>TG 2 (Tech. rep. #1)</th>
<th>TG 3 (Tech. rep. #1)</th>
<th>VG 1 (Tech. rep. #1)</th>
<th>VG 2 (Tech. rep. #1)</th>
<th>VG 3 (Tech. rep. #1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Absolute Control</td>
<td>Normal Control</td>
<td>Normal Control</td>
<td>Normal Control</td>
<td>Positive Control</td>
<td>Positive Control</td>
</tr>
</tbody>
</table>

Plate is treated with appropriate fluorescent probe

Plate is taken to the Plate reader for analysis
B

**Figure 2.3:** A: (Depicted above on pg 50) Example of an experimental layout used in plate reader studies. B: Summary of steps followed for each plate reader experiment.
2.3.2.1 NO Measurements – DAF

DAF-2/DA is a NO-specific fluorescent probe (Strijdom, Muller, & Lochner, 2004), which measures intracellular NO-production. Upon reacting with NO, DAF-2/DA is oxidized and emits a green fluorescence, which can be measured by the microplate reader. For cell treatment, DAF-2/DA was diluted with PBS to a final concentration of 10 µM, and then applied to the cells in the 24-well plate for 2 hours at 37 °C (Fig 2.4). After 2 hours, the DAF-2/DA-PBS supernatant was aspirated and the cells were washed once with PBS, and then covered in 500 µl PBS for microplate reader analysis.

Positive Control: DEA/NO

DEA/NO is a NO donor, and has previously been shown in our laboratory to be a consistent and reliable positive control for intracellular NO measured by DAF-2/DA. Designated positive control (+C) wells were first incubated with DAF-2/DA for 1.5 hours, before 100 µM DEA/NO was added for an additional period of 30 minutes. In other words, DEA/NO was added to the +C wells for the final 30 minutes of the 2-hour DAF-2/DA protocol (Fig 2.4 and 2.5).

Figure 2.4: A schematic representation of the protocol used for treatment with DAF-2/DA and positive NO control, DEA/NO.
Figure 2.5: DEA/NO (100 µM; 30 minutes) significantly increased mean DAF-2/DA fluorescence intensity compared to the normal control in non-serum starved samples and serum starved (SS) samples. DEA/NO was included as a positive NO control in all experiments (n = 6/ group). All values calculated as mean+SEM.
2.3.2.2 RNS Measurements – DHR

DHR-123 is an uncharged and non-fluorescent indicator of RNS (used as a marker of peroxynitrite in this study) (Tiede, Cook, Morsey, & Fox, 2012) that passively diffuse across the membrane, and is oxidized to form cationic rhodamine-123. Rhodamine-123 eventually localizes in the mitochondria and emits a green fluorescence that is detected by the microplate reader. For cell treatment, 2 µM DHR-123 was applied to the cells in the 24-well plate for 3 hours at 37 °C (Fig 2.6). The fluorescent probe was then washed out and further prepared as described in section 2.3.2.1.

Positive Control: Authentic peroxynitrite

Designated +C wells were first incubated with DHR-123 for 1 hour, before 100 µM peroxynitrite was added for an additional period of 2 hours. In other words, peroxynitrite was added to the +C wells for the final 2 hours of the 3-hour DHR-123 protocol (Fig 2.6 and 2.7).

![Figure 2.6](https://scholar.sun.ac.za)

**Figure 2.6:** A schematic representation of the protocol used for treatment with DHR-123 and positive control, authentic peroxynitrite.
Cell viability was assessed through the measurement of necrosis. During necrosis, there is a loss in cell membrane integrity. PI is able to enter the nucleus following the cell’s loss in membrane integrity and bind to the cell’s DNA (Riccardi & Nicoletti, 2006), allowing for the measurement of necrosis (Wilkins, Kutzner, Truong, Sanchez-Dardon, & McLean, 2002). For cell treatment, PI was diluted with PBS to a final concentration of 5 µM, and then applied to the cells in the 24-well plate for 15 minutes at 37 °C. After the 15-minute incubation period, PI was left on the plate and the cells were taken to the microplate reader for analysis.
Positive Control: dH$_2$O

When dH$_2$O is added to cells, it diffuses across a semi-permeable membrane from an area with low osmolarity (high H$_2$O concentration) to an area of high osmolarity (low H$_2$O concentration). Therefore the net result of treating cells with dH$_2$O is the movement of H$_2$O molecules over the cell membrane and into the cell. This subjects the cell to osmotic stress, which causes the cell to swell and eventually burst. Rupturing of the cell membrane due to necrosis allows the PI fluorescent probe to enter and stain the nucleus by intercalating with the DNA (Fig 2.8). Designated +C wells were treated with PI that was diluted in dH$_2$O, while all the other experimental group wells were treated with PI that was diluted in PBS.

![Figure 2.8](https://scholar.sun.ac.za)

**Figure 2.8:** dH$_2$O significantly increased PI staining compared to the normal control in non-serum starved samples and serum starved samples. dH$_2$O was included as a positive control in all experiments (n = 7/group). All values calculated as mean+SEM.
2.4 Signalling Investigations – Western Blot Analyses

AEC western blot experiments consisted of the following groups:

- Control (Contains Combined Vehicle Only)
- 1 nM Melatonin + Combined Vehicle
- High concentration ART + Combined Vehicle
- ART + Melatonin + Combined Vehicle

Unlike the microplate reader experimental groups, western blot experimental groups all contained a combined vehicle. This allowed for all groups to be directly compared without considering a vehicle effect. The reason for this alteration was due to the space restrictions associated with western blot sample loading. Limited loading space meant we had to reduce the number of experimental groups in order to cater for an adequate sample size (n = 5 / group). The consequence of this alteration was that all melatonin and ART concentrations had to be re-calculated. Refer to Appendix C for detailed melatonin, ARV and vehicle concentration calculations. A protein concentration of approximately 4.62 – 7.11 µg/µl was obtained from each petri dish.

2.4.1 Materials

- **Cell Signaling Technologies** (Beverly, MA, USA):  
  - eNOS  
  - phospho-eNOS (Ser 1177)  
  - Cleaved caspase-3  
  - Anti-rabbit immunoglobulin G, HRP- conjugated secondary antibody
- **Santa Cruz Biotechnologies** (Santa Cruz, CA, USA):  
  - Nitrotyrosine  
  - p22 PHOX  
  - iNOS
- Clarity™ enhanced chemiluminescence (ECL) detection reagent: *Bio-Rad* (Hercules, CA, USA)
- Primary and secondary SingalBoost™ Immunoreaction Enhancer: *Millipore* (Billerica, MA, USA).
- PageRuler™ Prestained Protein Ladder: *Thermo Scientific* (Lithuania, EU)
- All other chemicals and buffer reagents: *Sigma-Aldrich* (St Louis, Mo, USA) and *Merck* (Darmstadt, Germany).
2.4.2 Methods

The Western blotting procedure requires a high yield of protein. On the third passage, cells were re-seeded from small 35 mm petri dishes to large 100 mm petri dishes. A single large 100 mm petri dish was found to yield a sufficient protein concentration for one lysate.

2.4.2.1 Cell Lysates

After treatment, AECs were covered in 10 ml PBS and a rubber cell-scraper was used to gently lift the cells from the plate. The cell-PBS suspension was then transferred to the relevant tube and centrifuged for 3 minutes at 1000 rpm, after which the cell pellets were transferred to an eppendorff tube. Zirconium oxide beads (0.15 mm) were added to the pellet, along with 650 ul lysis buffer (Table 2.2) followed by homogenization with a Bullet Blender™ (Next Advance, Inc., NY, USA). 3 cycles of 1 minute were performed on setting 5 of the Bullet Blender. In between cycles, samples were allowed to rest for 5 minutes. Samples were placed on ice for 30 minutes after which they were centrifuged for 20 minutes at 15 000 rpm at 4 °C. Protein content of the supernatant was determined using the Bradford assay (Bradford, 1976). Based on the Bradford assay findings, samples were prepared containing 2X Laemmli buffer (Table 2.3), lysis buffer and protein lysate, where a final protein content of 40 μg/15 μl of sample was achieved.
Table 2.2: Lysis buffer contents and preparation.

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Final concentration in buffer</th>
<th>Amount of 10X stock for 30 ml</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris-HCl (pH 7.5)</td>
<td>20 mM</td>
<td>3.0 ml</td>
</tr>
<tr>
<td>EGTA</td>
<td>1 mM</td>
<td></td>
</tr>
<tr>
<td>EDTA</td>
<td>1.0 mM</td>
<td>300 µl</td>
</tr>
<tr>
<td>NaCl</td>
<td>150 mM</td>
<td>4.5 ml</td>
</tr>
<tr>
<td>β-glycerophosphate</td>
<td>1.0 mM</td>
<td>0.006 g</td>
</tr>
<tr>
<td>Tetra-sodium pyrophosphate</td>
<td>2.5 mM</td>
<td>0.03 g</td>
</tr>
<tr>
<td>NaF</td>
<td>50 nM</td>
<td>0.0639 g</td>
</tr>
<tr>
<td>Na₃VO₄</td>
<td>1.0 mM</td>
<td>3.0 ml</td>
</tr>
<tr>
<td>Triton-X100 (1 ml T in 10 ml dH₂O)</td>
<td>1.0 %</td>
<td>3.0 ml</td>
</tr>
<tr>
<td>Leupeptin</td>
<td>10 µg / ml</td>
<td>30 µl</td>
</tr>
<tr>
<td>Aprotonin</td>
<td>10 µg / ml</td>
<td>30 µl</td>
</tr>
<tr>
<td>Sodium dodecyl sylphate (SDS)</td>
<td>0.1 %</td>
<td>30 µl</td>
</tr>
<tr>
<td>PMSF</td>
<td>50 µg / ml</td>
<td>90 µl</td>
</tr>
<tr>
<td>Distilled water</td>
<td></td>
<td>Fill up to 30 ml</td>
</tr>
</tbody>
</table>

Table 2.3: Contents of 2X Laemmli buffer (Laemmli, 1970).

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>SDS</td>
<td>4 %</td>
</tr>
<tr>
<td>Glycerol</td>
<td>20 %</td>
</tr>
<tr>
<td>2-Mercaptoethanol</td>
<td>10%</td>
</tr>
<tr>
<td>Bromophenol blue</td>
<td>0.004 %</td>
</tr>
<tr>
<td>Tris HCl</td>
<td>0.125 M</td>
</tr>
</tbody>
</table>
2.4.2.2 Protein Loading and Transfer

Prepared lysates were boiled for 5 minutes before loading. A PageRuler™ ladder was loaded into the first well of every gel (7 µl) followed by 15 µl of lysate for each sample. 26-well (4–15% gradient) Criterion™ TGX Stain-Free™ Protein Gels (Bio-Rad, CA, USA) were used for all western blot experiments. Gels were run at 140 V for 10 minutes, followed by a second 40-minute run at 200 V. After running, the gels were activated for 1 minute using the Chemidoc™ MP Imager System with Image Lab™- 5 software (Bio-Rad, CA, USA), where the gels could be checked for equal protein loading. Following gel activation, proteins on the gel were transferred to a Trans-Blot® Turbo™ Midi PVDF membrane (Bio-Rad, CA, USA), using the Trans-Blot® Turbo™ Transfer System (Bio-Rad, CA, USA). The “Standard SD” protocol was used for all transfers and involved a 30 minute transfer at a constant voltage of 25 V and a varying current of up to 1 A. After the transfer the membrane was imaged on the Chemidoc™ again to ensure all proteins transferred correctly and evenly.

2.4.2.3 Immunodetection

Membranes were blocked for non-specific binding for 1.5 hours in 5 % fat-free milk in Tris-buffered saline, 0.1 % tween-20 (TBS-Tween). After blocking, membranes were washed for 30 minutes (3 x 10 minute cycles) with TBS-tween and then incubated overnight at 4 ºC with a specific primary antibody (Table 2.4). The next morning, the membranes were washed again for 30 minutes in TBS-tween and the incubated with an anti-rabbit immunoglobulin G, horseradish peroxidase (HRP) - conjugated secondary antibody for 1 hour at room temperature. The specific primary and secondary antibody conditions of each antibody are shown in Table 2.4. After the 1-hour incubation, membranes were rinsed again for 30 minutes, after which protein visualization took place. To visualize the proteins, membranes were incubated in Clarity™ for 5 minutes, after which the stain-free membranes were exposed on the Chemidoc™. Exposure time and the number of images captured were set beforehand and could be visualized throughout the procedure.

Image Lab™- 5 software standardises the visualised protein bands against the total amount of protein per lane transferred to the membrane (the image that was taken following transfer of protein to the membrane). The raw values generated by the software are then used to normalise all values to the control group, where the control is represented by the value 1. See Appendix D for the method used to normalise the data.
Some membranes were stripped of their original probe and re-probed for a different antibody. To strip membranes of primary and secondary antibodies, membranes were washed for 14 minutes (2 x 7 minute cycles) with dH₂O, then for 7 minutes with 0.2 M NaOH, followed by 14 minutes with dH₂O again. Membranes were then blocked in 5 % milk for 2 hours and could then be re-probed at a different molecular weight on the membrane than the initial antibody; therefore some proteins shared normalization membrane images.

**Table 2.4:** Specifications for each antibody after optimisation for Western blotting.

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Primary Antibody Dilution</th>
<th>Secondary Antibody Dilution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phospho-eNOS 140 kDa</td>
<td>1:1000 TBS (5µl in 5ml)</td>
<td>1:4000 5% Milk (1.25µl in 5ml)</td>
</tr>
<tr>
<td>Total-eNOS 140 kDa</td>
<td>1:1000 TBS (5µl in 5ml)</td>
<td>1:4000 5% Milk (1.25µl in 5ml)</td>
</tr>
<tr>
<td>iNOS 131 kDa</td>
<td>1:200 2.5% Milk (25µl in 5ml)</td>
<td>1:4000 5% Milk (1.25µl in 5ml)</td>
</tr>
<tr>
<td>Nitrotyrosine 90 kDa</td>
<td>1:5000 TBS (1µl in 5ml)</td>
<td>1:4000 TBS (1.25µl in 5ml)</td>
</tr>
<tr>
<td>p22 PHOX 22 kDa</td>
<td>1:500 1% Milk (10µl in 5ml)</td>
<td>1:4000 5% Milk (1.25µl in 5ml)</td>
</tr>
<tr>
<td>Cleaved Caspase-3 17/19 kDa</td>
<td>1:500 1° SignalBoost™ (10µl in 5ml)</td>
<td>1:4000 2° SignalBoost™ (1.25µl in 5ml)</td>
</tr>
</tbody>
</table>
2.5 ORAC Assay

The oxygen radical absorbance capacity (ORAC) assay is a method to assess the antioxidant capacity of biological samples (Huang, Ou, Hampsch-Woodill, Flanagan, & Prior, 2002; Prior et al., 2003). ORAC assays were kindly conducted by Dr D Blackhurst (Division of Chemical Pathology, University of Cape Town, SA).

AECs were passaged and maintained with the same method previously described in the section 2.4.2, and treated as previously described under the heading of section 2.4 (See Appendix C). After treatment, cells were washed with PBS and then covered with just over 1 ml of PBS. The cells were removed from the petri dish surface using a rubber cell scraper after which exactly 1 ml cell-PBS suspension was transferred to the relevant eppendorff tube. Sample tubes were then frozen and transported to the University of Cape Town (UCT) on dry-ice, for analysis.

For analysis, the cell samples were subjected to sonication using a Virtis VirSonic 100 (United Scientific (Pty) Ltd, CPT, SA) with 20 x 1 second bursts at 3 watts. At this stage 25 µl were removed for protein estimation by the Markwell modification of the Lowry method (Markwell, Haas, Bieber, & Tolbert, 1978), using bovine serum albumin as the standard. Cell samples were then centrifuged at 1000 x g at 4 °C for 15 minutes and the supernatant was used for ORAC measurement, using the method previously described by Maarman et al. (2015).

2.6 Statistical Analyses

All microplate reader data was calculated as mean ± standard error of the mean (SEM), with values expressed as a % of the control (control adjusted to 100%). For Western blot data, controls were adjusted to the value of 1. Total protein expression was calculated as a ratio of the loading control. For the ORAC assays, the antioxidant capacity is expressed in trolox equivalents. Trolox is a vitamin-E analogue that is used as the standard measure of antioxidant capacity.

One-way ANOVA with a Bonferroni post-hoc multiple comparison test were used to determine significance. Differences with a p-value < 0.05 were considered statistically significant. All data was analysed using GraphPad Prism 6 software (GraphPad Software, San Diego, CA, USA).

Sample sizes are indicated below each graph in Chapter 4.
3 - Materials and Methods: *Ex vivo* and *In vivo* studies

This chapter contains a description of the materials and methods used for the *ex vivo* and *in vivo* studies. The *ex vivo* studies consisted of investigations into the organ bath-based vascular contraction-relaxation function of aortas obtained from healthy male Wistar rats. In these studies, melatonin and/or ART were administered directly to the aortic rings mounted in the organ bath, and acute vascular and endothelium-dependent responses were measured. The *in vivo* studies comprised of an 8-week period of oral treatment with melatonin and/or ART, followed by organ bath-based investigations into the endothelium-dependent and endothelium-independent vascular contraction-relaxation function of the aortas obtained from the treated animals. The results for these studies can be found in Chapter 5.

### 3.1 Materials

- Anaesthetic (Sodium pentobarbitone; C₁₁H₁₇N₂NaO₃): *Bayer* (Isando, Gauteng, SA).
- *Sigma-Aldrich* (St Louis, Mo, USA):
  - Acetylcholine
  - Phenylephrine
  - Nω-Nitro-L-arginine methyl ester (L-NAME) hydrochloride
  - Sodium Nitroprusside (SNP)
- All other chemicals and buffer reagents: *Sigma-Aldrich* (St Louis, Mo, USA) and *Merck* (Darmstadt, Germany).

### 3.2 Ethics Clearance and Animal Care

All experiments were conducted in accordance with the accepted standards for the use of animals in research as set out by the South African National Standards document (SANS 10386: 2008) of the South African Bureau of Standards (SABS). Ethics approval was received from The Research Ethics Committee: Animal Care and Use of the University of Stellenbosch (US; Faculty of Medicine and Health Sciences; Protocol Number: SU-ACUD14-00021).
Age matched male Wistar rats with an initial weight range between 180 – 200 g were provided, housed, and cared for by the Animal Housing Unit of the Faculty of Medicine and Health Sciences of US. The animals were housed (2 – 3 rats per cage) in a temperature (22 °C) and humidity (40 %) controlled environment where they were subjected to a normal 12 hour artificial day / night cycle. Animals had free access to a standard rat chow diet as well as drinking water.

3.3 Experimental Protocols and Study Design

For aortic ring experiments, animals were transferred from the Animal Housing Unit to the animal care facility of the Division of Medical Physiology, 1 day prior to sacrifice (usually between 2 – 4 animals). This was to ensure animals had a day to recover from the stress of being transported as well as time to adapt to their new environment.

The animals were anesthetised through intraperitoneal (IP) injection of 160 mg / kg sodium pentobarbitone in the lower right abdomen. Deep anesthesia was confirmed by the disappearance of the pedal pain withdrawal reflex and absence of eye reflexes. Euthanasia was achieved by means of a clamshell thoracotomy and eventually exsanguination.

3.3.1 Excision and Mounting of Aortic Rings

Following the establishment of deep anesthesia, an incision was made through the skin and muscle layers across the ventral side of the rat, just below the thoracic region. The diaphragm and ribcage were cut in a cranial direction as to expose the thoracic cavity. The heart, lungs, trachea and oesophagus were removed. The thoracic aorta (above the diaphragm to distal end of the aortic arch) was excised and immediately placed in ice-cold Krebs Henseleit buffer (KHB) (Table 3.1). Using a magnifying glass, the perivascular fat and connective tissue was removed (Fig 3.1) and the aorta was cut into a 3 - 4 mm ring segment that was subsequently mounted onto two stainless steel hooks (Fig 3.2) in a 25 ml organ bath (AD Instruments, Bella Vista, NSW, Australia) containing oxygenated (95% O₂ and 5% CO₂) KHB. Aortic ring tension was recorded with an isometric force transducer (TRI202PAD, Panlab, lCornellà, BCN, Spain) and data was analysed with LabChart 7 software (Dunedin, New Zealand) (Westcott, 2015).
Table 3.1: Composition of Krebs Henseleit buffer.

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Concentration (mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NaCl</td>
<td>119</td>
</tr>
<tr>
<td>NaHCO₃</td>
<td>25</td>
</tr>
<tr>
<td>KCL</td>
<td>4.75</td>
</tr>
<tr>
<td>KH₂PO₄</td>
<td>1.2</td>
</tr>
<tr>
<td>MgSO₄.7H₂O</td>
<td>0.6</td>
</tr>
<tr>
<td>Na₂SO₄</td>
<td>0.6</td>
</tr>
<tr>
<td>CaCl₂.H₂O</td>
<td>1.25</td>
</tr>
<tr>
<td>Glucose</td>
<td>10</td>
</tr>
</tbody>
</table>

Figure 3.1: A thoracic aorta excised and cleaned of connective tissue and perivascular fat (Loubser, 2014).
3.3.2 Ex Vivo Studies

For all *ex vivo* aortic ring experiments, phenylephrine (Phe) was used to induce contraction, and acetylcholine (Ach) was used to induce relaxation. Phe is an α-adrenergic receptor agonist which acts directly on the VSMCs eventually leading to contraction. Ach binds to endothelial surface receptors (Fig 1.3) resulting in an increase in intracellular calcium and consequently eNOS activation. eNOS activation leads to the release of NO which diffuses to the VSMC layer resulting in VSMC relaxation.

The isometric tension measurement protocol used was a modified version of a technique previously described by Privet *et al.* (2004). The mounted aortic rings were stabilised for 30 minutes at a resting tension of 1.5 g, over which period, the KHB was changed every 10 minutes with fresh, pre-warmed (37 °C) KHB. After the 30-minute stabilisation, an initial functionality test was conducted, whereby 100 nM Phe was added to the organ bath. Once the ring was no longer contracting (a plateau was reached), 10 µM Ach was added. Rings that showed at least a 70 % relaxation of maximum phenylephrine-induced contraction during the initial functionality test were included for further investigations. Following the functionality test, the organ bath was rinsed 3 times with fresh, warm KHB and the rings were once again stabilised for 30 minutes at a tension of 1.5 g, where the buffer was changed every 10 minutes. After the second 30-minute stabilisation period, the relevant treatment was added to the organ bath, which was followed by a 30-minute incubation period.
period. Following drug administration and the 30-minutes incubation, aortic ring contraction was induced with the cumulative administration of Phe (100 nM – 1 µM). Once a plateau was reached, aortic rings were subsequently exposed to cumulative concentrations of Ach (30 nM – 10 µM). Experiments were terminated once the aortic rings experienced maximum relaxation. Figure 3.3 is a representative LabChart recording showing the aortic ring responses to the experimental protocol followed for the ex vivo studies.

Ex vivo aortic ring experiments consisted of the following groups:

- Control (Contains Combined Vehicle Only)
- 10 µM Melatonin (Sigma Aldrich, St Louis, MO, USA) + Combined Vehicle
- High concentration ART (SantaCruz Biotechnology, CA, USA) + Combined Vehicle
- ART + Melatonin + Combined Vehicle

Ex vivo treatment groups all contained a combined vehicle. This allowed for all groups to be directly compared without considering a vehicle effect. A very limited number of rats were allocated to the ex vivo portion of this study due to the adjusted protocol’s novel and experimental nature. A limited number of rats meant we had to reduce the number of experimental groups in order to cater for an adequate sample size (n = 5 - 6). For ex vivo studies, a melatonin concentration of 10 µM was used, unlike the 1 nM melatonin concentration, which was used for the in vitro studies. The reason for this discrepancy is that the ex vivo studies were conducted before the in vitro melatonin dose-response experiments were completed, and at that stage, the assumption was made that the higher dose of melatonin would be more effective. It only became evident that the lower dose of melatonin was in fact the more effective concentration to use, once melatonin dose-response experiments were completed. Refer to Appendix E for detailed ex vivo melatonin, ARV and vehicle concentration calculations.
For the in vivo studies, 80 male Wistar rats were divided into the following 4 experimental groups (Fig 3.4):

- Control (Untreated)
- Melatonin (Sigma Aldrich, St Louis, MO, USA)
- ART/Odimune® (Cipla MedPro, Bellville, WC, SA)
- ART + Melatonin

One week prior to the start of treatment, rats weighing 180 – 200g were randomly separated into the 4 treatment groups. Between 2 and 3 rats were housed per cage according to the conditions mentioned in section 3.2. During this week, the water-intake of the rats was measured and recorded, but the rats received no other treatment.
3.3.3.1 Drug Administration

Following the initial 1-week water-intake monitoring, an 8-week treatment period was initiated. Throughout the 8-week treatment period, rats were weighed once per week (Monday morning) and all fluid intake was monitored and measured on a daily basis. Rats were caged together in groups of 2 or 3, and therefore the amount of fluid drank per cage was measured, and then divided based on the number of rats per cage.

10 mg/kg melatonin was found to be effective in reducing oxidative damage (Cardinali, Cano, Jimenez-Ortega, & Esquifino, 2011; Favero et al., 2017), and is the current concentration of melatonin used in other studies within our laboratory. This dose of melatonin was administered through the drinking water, where each cage containing a given number of rats was given 200 ml a day, and the melatonin-water was measured the next day at the same time, to assess the amount of water consumed by each cage daily. This amount was then divided by the amount of rats in the given cage in order to calculate the mean water consumed by each rat within a single cage. This information, along with the average weight of the rats in a specific cage allowed for the calculation of the amount of melatonin each cage received for a specific week. This was necessary to ensure the rats were always receiving the prescribed dose of melatonin. For example, in cages that consumed very high quantities of melatonin-water, a lower stock concentration of melatonin was prepared, while cages that drank very little water, needed higher concentrations of melatonin in their drinking water to compensate for their reduced water consumption. Melatonin powder was dissolved in methanol and made up in stocks every
two days, while the drinking water containing melatonin was replaced every day. The amount of methanol used to dissolve the melatonin powder was dependent on the amount of melatonin stock that was made up, where 0.2% of the amount of stock being made was used to determine the quantity of methanol used. For example, if a 500 ml melatonin stock was prepared, 1 ml of methanol (0.2% x 500 ml) was used. All drinking bottles containing melatonin were covered with two layers of aluminium foil and a metal casing to ensure the melatonin was not inactivated by light. For detailed melatonin calculations, stock preparation and administration see Appendix F.

For ART treatment, an ART-water drug suspension was administered daily by oral gavaging by a qualified staff member (Mr. Noël Markgraaff; Manager: Care and Use of Laboratory Animals, SU). Groups that did not receive ART were also gavaged, but with water instead of the ART-suspension. The procedure of gavaging involved the insertion of a feeding tube along the roof of the animal’s oral cavity towards the animal’s left side and down the animal’s esophagus and into the stomach of the animal. The ART-suspension or water was therefore injected directly in the stomach of the animal. The rats received 1 ml ART-suspension or water per day. The ART-suspension was made up weekly, according to the average weekly weight of the cage. To make up the ART-suspension, Odimune® tablets were ground into a fine powder using a pestle and mortar. The appropriate amount of Odimune® powder was then weighed and added to dH₂O, after which the mixture was thoroughly mixed and stored in the refrigerator.

Each Odimune® tablet contains the human recommended daily dose of active ingredients (EFV: 600 mg; FTC: 200 mg; TDF: 300 mg) for an average person (70 kg). These values, as well as the weekly rat weights, were used to calculate how much tablet powder needed to be combined with dH₂O in order for the rats to receive the applicable concentrations of the three individual ARVs (EFV: 51.6 mg/kg; FTC: 17.4 mg/kg; TDF: 25.8 mg/kg). See Appendix G for the method used to calculate the weekly drug dose.

### 3.3.3.2 Aortic Ring Investigations

The thoracic aorta was excised, cleaned of connective tissue and suspended in the organ bath as described under section 3.3.1. For in vivo experiments, two types of aortic ring investigations were conducted. For the first set of experiments we conducted endothelium-dependent relaxation studies, were Phe was used to induce contraction and Ach was used
to induce relaxation (as previously described in section 3.3.2). As a control for these endothelium-dependent relaxation studies, some of the rings in this section were pre-treated with the NOS inhibitor, L-NAME (100 μM) 15 minutes prior to the administration of cumulative amounts of Phe and Ach. For these experiments, a single aorta was cut into two rings, which were mounted immediately in separate organ baths (Fig 3.5). The normal Phe-Ach experiments were always conducted in Organ Bath 1, which L-NAME studies were always conducted in Organ Bath 2.

For the second set of experiments we conducted endothelium-independent relaxation studies, where Phe was once again used to induce contraction, however, for these studies sodium nitroprusside (SNP) - an exogenous NO donor - was used to induce relaxation. Since SNP is a NO donor, relaxation observed is independent of the endothelium, as the NO released from SNP diffuses directly into the VSCMs. For SNP investigations, a single aorta was once again cut into two rings (Fig 3.5), and all SNP studies were conducted in Organ Bath 2.

Figure 3.5 shows how aortas were separated and cut for the two different studies.
Figure 3.5: Scheme indicating procedures performed on cleaned aortic tissue to obtain aortic rings for isometric tension studies.
Following the initial stabilisation period, functionality test (done with Phe and Ach for all experiments), and the second stabilisation period (as described in section 3.3.2), each group was subjected to the following three isometric tension protocols:

1) **Cumulative Phe-induced contraction followed by cumulative Ach-induced relaxation:** Cumulative administration of Phe results in a step-wise increase in the total Phe concentration as follows: 100 nM; 300 nM; 500 nM; 800 nM; 1 µM. This administration results in cumulative aortic ring contractions. Each dose of Phe was administered directly to the organ bath as soon as maximum contraction was reached with the previous administration. Once the final dose of Phe was administered to produce a final Phe concentration of 1 µM, maximum contraction was reached and therefore Ach was administered in order to induce endothelium-dependent relaxation. Ach was also administered in a cumulative manner resulting in step-wise increases in Ach concentrations: 30 nM; 100 nM; 300 nM; 1 µM; 10 µM. The experimental protocol was terminated once the final Ach administration (final concentration: 10 µM) resulted in maximum % relaxation of contraction (Fig 3.6).

**Figure 3.6:** A representative LabChart recording showing the aortic ring responses to the experimental protocol followed for Phe administration followed by Ach. This graph was from an aortic ring in the control group.
2) **L-NAME pre-treatment, followed by cumulative Phe-induced contraction and cumulative Ach-induced relaxation:** In this protocol, the role of NOS-derived NO was manipulated by pre-administration of the NOS-inhibitor, L-NAME (100 μM) 15 minutes prior to the cumulative Phe - Ach protocol (Fig 3.7).

![Diagram showing tension changes over time](image)

**Figure 3.7:** A representative LabChart recording showing the aortic ring responses to the experimental protocol followed for L-NAME pre-treatment followed by Phe and Ach administration. This graph is from an aortic ring in the control group.
3) **Cumulative Phe-induced contraction followed by cumulative SNP-induced relaxation:** Phe was administered in a cumulative fashion as mentioned in the first protocol. Upon reaching maximum contraction, SNP was administered to induce endothelium-independent relaxation. SNP was also administered in a cumulative manner resulting in step-wise increases in SNP concentrations: 1.2 nM; 20 nM; 70 nM; 100 nM; 750 nM. The experimental protocol was terminated once the final SNP administration (final concentration: 750 nM) resulted in maximum % relaxation of contraction (Fig 3.8). SNP concentrations used were based on various sources in the literature (Ajay & Mustafa, 2006; Ameer, Boyd, Butlin, Avolio, & Phillips, 2015; Banda, Lefer, & Granger, 1997; Bonaventura, Lunardi, Rodrigues, Neto, & Bendhack, 2008; Sutliff et al., 2002). See **Appendix H** for detailed SNP concentration calculations and administration.

**Figure 3.8:** A representative LabChart recording showing the aortic ring responses to the experimental protocol followed for Phe administration followed by SNP. This graph is from an aortic ring in the control group.
3.4 Signalling Investigations – Western Blot Analyses

3.4.1 Materials

The same materials were used as described in Chapter 2, section 2.4.1

3.4.2 Methods

Due to the fact that for most samples, only a very small piece of aorta was frozen away, two samples from the same experimental group were combined to make one lysate.

3.4.2.1 Aortic Tissue Lysates

Aortic tissue that had been previously frozen at -80 °C was stored in liquid nitrogen, along with pulverizing equipment and any tools used to scrape or transfer the frozen aortic tissue. All these items were kept cold in liquid nitrogen so that when the tissue was pulverized it formed a fine powder. Following aortic tissue pulverization, 60 – 106 mg of the aortic tissue-powder was placed in an eppendorff tube along with the equivalent mass of 1.6 mm stainless steel beads and 600 µl complete lysis buffer (Table 2.2). Samples were then homogenised in a Bullet Blender™ (Next Advance, Inc., NY, USA) with the following protocol:

- 3 minutes at speed 8
- 5 minutes rest
- 2 minutes at speed 10

Samples were placed on ice for 30 minutes after which they were centrifuged for 20 minutes at 15 000 rpm at 4 °C. Protein content was determined by the Bradford assay (Bradford, 1976) as mentioned in Chapter 2, section 2.4.2.1. A final protein content of 60 µg/15 µl of sample was achieved.

3.4.2.2 Protein Loading, Transfer and Measurement

Loading of samples onto the Criterion™ TGX Stain-Free™ Protein Gels, subsequent transfer to membrane as well as antibody conditions and protein band visualization was conducted as described in Chapter 2, sections 2.4.2.2 and 2.4.2.3.

3.5 ORAC Assay

ORAC assays were kindly conducted by Dr D Blackhurst (Division of Chemical Pathology, University of Cape Town, SA).
Aortic tissue for the ORAC assay was prepared through pulverisation in the same manner described above in section 3.4.2.1. Aortic tissue-power was placed in an empty eppendorff tube where the amount of tissue in mg within each sample tube was recorded. After pulverisation, samples were immediately frozen in liquid nitrogen and then transported to UCT on dry-ice, for analysis.

For analysis, aortic samples were homogenised (using an in-house ‘glass-ball’) and subjected to the same sonication, protein estimation and ORAC measurement protocols described in chapter 2, section 2.5.

### 3.6 Statistical Analyses

Data were analysed using GraphPad Prism 6 software (GraphPad Software, San Diego, CA, USA). Aortic ring contraction (Phe) studies are expressed as an increase in gram tension from a resting tension of 1.5 g, while relaxation (Ach/SNP) studies are expressed as % relaxation of the maximum contraction. Data were statistically analysed by means of a two-way ANOVA with a Bonferroni post-hoc multiple comparison test. Additionally, \( E_{\max} \) values were compared using a one-way ANOVA with a Bonferroni post-hoc multiple comparison test. Differences with a p-value < 0.05 were considered statistically significant.

For Western blot data, controls were adjusted to the value of 1. Total protein expression was calculated as a ratio of the loading control.

For the ORAC assays, the antioxidant capacity is expressed in trolox equivalents.

For western blot and ORAC data, one-way ANOVA with a Bonferroni post-hoc multiple comparison test were used to determine statistical significance. Differences with a p-value < 0.05 were considered statistically significant.

Sample sizes are indicated below each graph in Chapter 5.
4 – Results: In vitro studies

Chapter 4 describes and reports data collected from the in vitro study, the materials and methods for which can be found in Chapter 2.

4.1 Plate Reader Analyses

The initial aim of the in vitro study set out to determine the individual optimum concentrations of both melatonin and first line FDC ART treated aortic endothelial cells (AECs) at which the greatest differences were seen when compared to untreated AECs, with respect to NO production, RNS production and cell viability. This was investigated through dose-response studies whereby AECs were treated with various melatonin and ART concentrations for a fixed period.

For the second aim of the in vitro study, the optimum concentrations obtained in the dose-response experiments above were used to treat AECs with both melatonin and ART in combination, in order to analyse any potential synergistic/antagonistic interactions between the two with respect to, NO production, RNS production and cell viability. For these studies, a single melatonin and FDC ART concentration was chosen for cell treatment over a period of 24 hours.

Following drug treatment, cells were treated with the applicable fluorescent probes, and the consequent cell fluorescence was measured using a plate reader. All treatment groups were compared to an untreated control group. Individual treatment groups were also compared to their respective vehicles. If no significant difference was seen between a group and its respective vehicle, then any significant changes between that particular treatment group and the control group were considered a vehicle effect. The n value represents the number of individual experiments (biological replicates) that were conducted for each endpoint, where each treatment group within an experiment consisted of 2 to 3 samples (technical replicates) for which the mean value was calculated. All values expressed as a % of the control (control adjusted to 100%). A p-value of < 0.05 was considered statistically significant.

4.1.1 Melatonin Dose-Response Studies

Dose-response experiments comprised of a 24-hour serum starvation (SS) period, followed by a 24-hour melatonin treatment period, followed by probe treatment and plate reader analyses. As previously mentioned in section 2.2, serum starvation (SS) was...
introduced to induce cell cycle synchronization, which enhances the cells’ responses to experimental treatment (Gerber et al., 1998; Russell & Hamilton, 2014), in otherwise healthy cells. Three treatment groups for melatonin were included: 1 nM; 1 µM and 10 µM melatonin. Results from these experiments (24-hour SS followed by 24-hour melatonin treatment) showed few to no changes between treatments groups, we therefore decided to introduce a 1-hour melatonin treatment period following 24-hour SS, to assess the early effects of melatonin on the cells.

4.1.1.1 24h SS + 24h Melatonin Treatment

DAF-2/DA: NO Production

AECs exposed to 24 hours of SS followed by treatment with 1 nM (103.0 ± 2.302%), 1 µM (103.3 ± 3.432%) and 10 µM (101.8 ± 3.544%) melatonin for 24 hours, showed no differences in mean DAF-2/DA fluorescence (NO production) compared to the untreated control group (100.0 ± 1.662%) (Fig 4.1). No changes were observed in any vehicle groups.

Figure 4.1: The effects of 24h SS followed by 24h melatonin treatment on NO production measured by DAF-2/DA fluorescence. NO production expressed as % DAF-2/DA stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.
DHR-123: RNS Production

AECs exposed to 24 hours of SS followed by treatment with 1 nM (109.3 ± 3.287%), 1 µM (104.6 ± 3.266%) and 10 µM (105.8 ± 2.737%) melatonin for 24 hours, showed no differences in mean DHR-123 fluorescence (RNS production) compared to the untreated control group (100.0 ± 1.080%) (Fig 4.2). No changes were observed in any vehicle groups.

Figure 4.2: The effects of 24h SS followed by 24h melatonin treatment on RNS production measured by DHR-123 fluorescence. RNS production expressed as % DHR-123 stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100%), n = 3/group.
PI: Cell Viability (Necrosis)

AECs exposed to 24 hours of SS followed by treatment with 1 nM (92.56 ± 3.114%) melatonin for 24 hours, showed a significant decrease in mean PI fluorescence (necrosis), when compared to the untreated control group (100.0 ± 1.041%), and its own vehicle (105.3 ± 0.9718%) (Fig 4.3). 1 µM (100.9 ± 1.703%) and 10 µM (102.3 ± 2.075%) melatonin treated groups showed no differences in mean PI fluorescence compared to the control group (100.0 ± 1.041%). No changes were observed in 1 µM and 10 µM vehicle groups.

* : p=0.0004 vs control
# : p=0.0004 vs 1nM vehicle

Figure 4.3: The effects of 24h SS followed by 24h melatonin treatment on necrosis measured by PI fluorescence. Necrosis expressed as % PI stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/group.
4.1.1.2 24h SS + 1h Melatonin Treatment

DAF-2/DA: NO Production

AECs exposed to 24 hours of SS followed by treatment with 1 nM (104.0 ± 1.238%), 1 µM (100.5 ± 0.7188%) and 10 µM (101.5 ± 0.8466%) melatonin for 1 hour, showed no differences in mean DAF-2/DA fluorescence (NO production) compared to the untreated control group (100.0 ± 0.9661%) (Fig 4.4). No changes were observed in any vehicle groups.

Figure 4.4: The effects of 24h SS followed by 1h melatonin treatment on NO production measured by DAF-2/DA fluorescence. NO production expressed as % DAF-2/DA stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.
DHR-123: RNS Production

AECs exposed to 24 hours of SS followed by treatment with 1 nM (101.2 ± 1.517%), 1 µM (98.5 ± 1.111%) and 10 µM (98.92 ± 1.196%) melatonin for 1 hour, showed no differences in mean DHR-123 fluorescence (RNS production) compared to the untreated control group (100.0 ± 0.9129%) (Fig 4.5). No changes were observed in any vehicle groups.

Figure 4.5: The effects of 24h SS followed by 1h melatonin treatment on RNS production measured by DHR-123 fluorescence. RNS production expressed as % DHR-123 stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 4/ group.
PI: Cell Viability (Necrosis)

AECs exposed to 24 hours of SS followed by treatment with 1 nM (81.89 ± 3.572%), 1 µM (86.89 ± 2.927%) and 10 µM (91.22 ± 2.133%) melatonin for 1 hour, showed a significant decrease in mean PI fluorescence (necrosis) respectively, when compared to the untreated control group (100.0 ± 0.7993%). All melatonin treatment groups also significantly decreased compared to their respective vehicles (1nM Veh: 96.11 ±1.844%; 1µM Veh: 94.89 ± 1.409%; 10µM Veh: 98.44 ± 1.042%). The 1 nM melatonin treatment group also showed a significant decrease in PI fluorescence compared to the 10 µM melatonin treatment group (Fig 4.6).

![Graph showing the effects of melatonin on PI fluorescence](image)

* : p<0.0001 vs control  
** : p<0.0001 vs control  
***: p<0.0001 vs control  
#: p<0.0001 vs 1 nM vehicle  
$: p<0.0001 vs 1 µM vehicle  
@: p<0.0001 vs 10 µM vehicle  
%: p<0.0001 vs 10 µM melatonin

**Figure 4.6:** The effects of 24h SS followed by 1h melatonin treatment on necrosis measured by PI fluorescence. Necrosis expressed as % PI stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.
4.1.2 ART Dose-Response Studies

Dose-response experiments comprised of a 24-hour serum starvation (SS) period, followed by a 24-hour ART treatment period, followed by probe treatment and plate reader analyses. Three treatment groups for ART were included: Low concentration ART (EFV: 5µM; FTC: 5µM; TDF: 80nM); Mid concentration ART (EFV: 8µM; FTC: 7.5µM; TDF: 400nM) and High concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM). In contrast to the melatonin dose-response studies, a 1-hour treatment period was not included for the ART dose-response studies. The shorter 1-hour treatment period was excluded for the ART studies since it would have little clinical relevance.
4.1.2.1 24h SS + 24h ART Treatment

DAF-2/DA: NO Production

AECs exposed to 24 hours of SS followed by treatment with low (100 ± 2.703% vs Veh: 108.6 ± 1.494%), mid (100.8 ± 3.116% vs Veh: 109.9 ± 2.394%) and high (112.9 ± 2.171% vs Veh: 105.4 ± 1.587%) concentration ART for 24 hours, all showed significant differences compared to their relative vehicles. However, only high concentration ART showed a significant increase in mean DAF-2/DA fluorescence (NO production) compared to the untreated control group (100.1 ± 2.224%). High concentration ART also increased NO production significantly compared to both low and mid concentration ART (Fig 4.7).

Figure 4.7: The effects of 24h SS followed by 24h ART treatment on NO production measured by DAF-2/DA fluorescence. NO production expressed as % DAF-2/DA stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.
DHR-123: RNS Production

AECs exposed to 24 hours of SS followed by treatment with low (108.6 ± 2.667%), mid (108.8 ± 2.783%) and high (108.8 ± 2.197%) concentration ART for 24 hours, all showed a significant increase in mean DHR-123 fluorescence (RNS production) compared to the untreated control group (100.0 ± 0.7896%). Only high concentration ART increased significantly compared to its vehicle (101.1 ± 2.539%) (Fig 4.8), meaning the increase in DHR-123 fluorescence seen in the low and mid concentration ART groups compared to the control were vehicle effects.

* EFV=5uM; FTC = 5uM; TDF= 80nM
** EFV=8uM; FTC = 7.5uM; TDF= 400nM
*** EFV=12uM; FTC = 10uM; TDF= 1uM

#: p=0.0152 vs control
###: p=0.0015 vs control
####: p=0.0015 vs control
$: p=0.0015 vs [high] vehicle

Figure 4.8: The effects of 24h SS followed by 24h ART treatment on RNS production measured by DHR-123 fluorescence. RNS production expressed as % DHR-123 stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.
PI: Cell Viability (Necrosis)

AECs exposed to 24 hours of SS followed by treatment with high (107.3 ± 1.344%) concentration ART for 24 hours, showed a significant increase in mean PI fluorescence (necrosis) compared to the untreated control group (100.0 ± 0.8571%). Low (101.3 ± 1.537%) and mid (101.7 ± 1.269%) concentration ART showed no differences in mean PI fluorescence when compared to the control. Only high concentration ART increased PI fluorescence significantly compared to its vehicle (101.3 ± 1.796%). High concentration ART also increased necrosis significantly compared to both the low and mid concentration ART groups (Fig 4.9).

Figure 4.9: The effects of 24h SS followed by 24h ART treatment on necrosis measured by PI fluorescence. Necrosis expressed as % PI stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/group.
4.1.3 Combination Studies

As previously mentioned in section 2.2.3, only a single concentration was used for both melatonin and ART in the combination studies. The individual melatonin and ART concentrations were selected based on the results observed in the dose-response experiments. Based on the above data, 1 nM melatonin and high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) were used in the combination studies. A 24-hour SS period was once again used, followed by a 24-hour treatment period.

DAF-2/DA: NO Production

AECs were exposed to 24 hours of SS followed by treatment with 1 nM melatonin (99.83 ± 1.515%), high concentration ART (98.00 ± 1.571%), as well as a combination of 1 nM melatonin + high concentration ART (99.33 ± 2.060%) for 24 hours. Results showed no differences in mean DAF-2/DA fluorescence (NO production) between the treatment groups and the untreated control group (100.0 ± 1.124%) (Fig 4.10). No changes were observed in any vehicle groups.

![Graph showing NO production](image)

*** EFV=12µM; FTC = 10µM; TDF= 1µM

**Figure 4.10:** The effects of 24h SS followed by 24h melatonin and ART combination treatment on NO production measured by DAF-2/DA fluorescence. NO production expressed as % DAF-2/DA stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.
DHR-123: RNS Production

AECs were exposed to 24 hours of SS followed by treatment with 1 nM melatonin (101.3 ± 0.9574%), high concentration ART (98.33 ± 0.8819%), as well as a combination of 1 nM melatonin + high concentration ART (98.89 ± 1.124%) for 24 hours. Results showed no differences in mean DHR-123 fluorescence (RNS production) between the treatment groups and the untreated control group (100.0 ± 1.124%) (Fig 4.11). No changes were observed in any vehicle groups.

**Figure 4.11:** The effects of 24h SS followed by 24h melatonin and ART combination treatment on DHR-123 fluorescence. RNS production expressed as % DHR-123 stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %). n = 3/ group.

*** EFV=12uM; FTC = 10uM; TDF = 1uM
**PI: Cell Viability (Necrosis)**

AECs were exposed to 24 hours of SS followed by treatment with 1 nM melatonin (92.43 ± 3.753%), high concentration ART (121.3 ± 9.114%), as well as a combination of 1 nM melatonin + high concentration ART (94.17 ± 5.082%) for 24 hours. When compared to the untreated control group (100.0 ± 1.067%), 1 nM melatonin was seen to significantly decrease mean PI fluorescence (necrosis). In contrast, the high concentration ART treatment group significantly increased mean PI fluorescence compared to the control, as well as compared to the 1 nM melatonin treatment group. When combined, the melatonin + ART treatment group significantly decreased mean PI fluorescence compared to the group treated with only ART. All treatment groups showed significant differences when compared to their relative vehicles (1nM Mel Veh: 111.2 ± 4.277%; High ART Veh: 97.83 ± 3.381%; Mel + ART Veh: 109.5 ± 3.413%) (Fig 4.12).

**Figure 4.12:** The effects of 24h SS followed by 24h melatonin and ART combination treatment on PI fluorescence. Necrosis expressed as % PI stained cells (calculated as a percentage of the untreated control group; Control fluorescence adjusted to 100 %), n = 3/ group.

*** EFV=12uM; FTC = 10uM; TDF= 1uM

*: p<0.0001 vs control  #: p<0.0001 vs 1 nM vehicle

**: p<0.0001 vs control  $: p<0.0001 vs [high] vehicle

@: p<0.0001 vs 1 nM melatonin

%: p<0.0001 vs melatonin + ART vehicle

&: p<0.0001 vs [high] ART
4.2 Western Blot Analyses

For *in vitro* cell signaling protein investigations, AECs were serum starved for 24 hours and then treated with a combination of 1 nM melatonin and high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) for a further 24 hours. The cells were used to make protein lysates as described in section 2.4. All western blot experimental groups contained a combined vehicle, which allowed for all groups to be directly compared without considering a vehicle effect.

For Western blot data, controls were adjusted to the value of 1. Total protein expression was calculated as a ratio of the loading control. A p-value of < 0.05 was considered statistically significant.

4.2.1 NO & RNS Signalling

For NO and reactive nitrogen species (RNS) signalling, total- and phosphorylated eNOS; total iNOS and total nitrotyrosine were examined. eNOS and iNOS are important signaling molecules in the production of NO, while nitrotyrosine is a marker of nitrosative stress.

4.2.1.1 eNOS

eNOS phosphorylation was significantly lower is the melatonin (0.6354 ± 0.04195), ART (0.4348 ± 0.05156) and melatonin + ART (0.2535 ± 0.03983) treatment groups compared to the control (1.000 ± 0.1068). The melatonin + ART treatment group also showed significantly less levels of eNOS phosphorylation compared to the melatonin treatment group (Fig 4.13 A).

No significant differences were seen in total eNOS expression between any of the treatment groups (Mel: 0.9188 ± 0.1187; ART: 0.7479 ± 0.09084; Mel + ART: 0.8895 ± 0.04375) when compared to the control (1.000 ± 0.1330) (Fig 4.13 B).

When eNOS phospho/total (P/T) ratios were calculated, the same pattern of significance emerged as was seen for the eNOS phosphorylation levels. All treatment groups showed significantly lower P/T eNOS levels (Mel: 0.5630 ± 0.01973; ART: 0.4298 ± 0.02480; Mel + ART: 0.2614 ± 0.05717) compared to the control (1.000 ± 0.1207), while the melatonin + ART treatment group had lower P/T eNOS levels when compared to the melatonin treatment group (Fig 4.13 C).
Figure 4.13: Changes in the eNOS phosphorylation and expression (Ser 1177) of AECs treated with melatonin and/or ART, for 24 hours. A) Analysed results and western blot for phospho-eNOS (Ser 1177); B) Analysed results and western blot for total-eNOS; C) Phosphorylated over total (P/T) ratio of eNOS (Ser 1177) (n=5/group).
4.2.1.2 iNOS

No band was detected for iNOS and therefore no statistical analyses could be conducted (Fig 4.14).

Figure 4.14: No iNOS band was detected in AECs treated with melatonin and/or ART, for 24 hours. Shown is the western blot for iNOS (n=5/group).

4.2.1.3 Nitrotyrosine

Nitrotyrosine levels in the ART treatment group (2.309 ± 0.3443) was significantly higher compared to the control group (1.000 ± 0.1707). When melatonin and ART were combined (1.266 ± 0.2876), nitrotyrosine levels decreased significantly compared to the group that was treated with ART alone (Fig 4.15). There were no significant differences seen in the melatonin group (2.825 ± 1.564).

Figure 4.15: Changes in nitrotyrosine levels of AECs treated with melatonin and/or ART, for 24 hours. Shown are the analysed results and western blot for nitrotyrosine (n=5/group).
4.2.2 ROS Signalling

For reactive oxygen species (ROS) signalling, total p22 PHOX was examined. p22 PHOX is a marker for NADPH-oxidase derived superoxide (Fukui et al., 1997), a reactive oxygen anion.

4.2.2.1 p22 PHOX

All treatment groups showed decreased p22 PHOX expression compared to the control (1.000 ± 0.1518). Both the ART (0.2039 ± 0.04277) and melatonin + ART (0.2587 ± 0.008877) combination treatment groups also showed significantly decreased p22 PHOX expression compared to the melatonin (0.5217 ± 0.08199) treatment group (Fig 4.16).

![Figure 4.16: Changes in the p22 PHOX expression of AECs treated with melatonin and/or ART, for 24 hours. Shown are the analysed results and western blot for p22 PHOX (n=5/group).](https://scholar.sun.ac.za)

4.2.3 Cell Viability Signalling

For cell viability signalling, cleaved caspase-3 was examined. Cleaved caspase-3 is a marker of apoptosis (Kuribayashi, Mayes, & El-Deiry, 2006), which is a programmed form of cell death.

4.2.3.1 Cleaved Caspase-3

The ART (0.2519 ± 0.03783) and melatonin + ART (0.3878 ± 0.07965) combination treatment groups showed significantly decreased cleaved Caspase-3 expression compared to the control (1.000 ± 0.1775), as well as compared to the melatonin treatment group (0.7586 ± 0.08004) (Fig 4.17).
for in vitro cell antioxidant capacity analyses, AECs were serum starved for 24 hours and then treated with a combination of 1 nM melatonin and high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) for a further 24 hours. All experimental groups contained a combined vehicle, which allowed for all groups to be directly compared without considering a vehicle effect.

Antioxidant capacity is expressed in trolox equivalents, which is a vitamin-E analogue that is used as the standard measure of antioxidant capacity (Maarman et al., 2015). A p-value of < 0.05 was considered statistically significant.

4.3.1 ORAC Assay

No significant differences were seen between melatonin (0.3324 ± 0.01852), ART (0.3134 ± 0.01466) and melatonin + ART (0.3366 ± 0.01324) treatment groups when compared to the control (0.2710 ± 0.04975) (Fig 4.18).

Figure 4.17: Changes in cleaved Caspase-3 expression of AECs treated with melatonin and/or ART, for 24 hours. Shown are the analysed results and western blot for cleaved Caspase-3 (n=5/group).
**Figure 4.18:** Antioxidant capacity of AECs treated with melatonin and ART, as well as a combined vehicle for 24 hours (n = 5/group).
5 – Results: *Ex vivo* and *In vivo* studies

Chapter 5 describes and reports data collected from the *ex vivo* and *in vivo* studies, the materials and methods for which can be found in Chapter 3. See Appendix I for biometric data obtained from the treated rats uduring the treatment protocol.

5.1 *Ex Vivo* Studies

The main aim of the *ex vivo* studies was to determine the acute effects of melatonin and ART administration on the vascular reactivity of untreated, control male Wistar rat aortas. To achieve this, drugs were administered as a pre-treatment, directly to an *ex vivo* organ bath containing a stabilised rat aortic ring. For *ex vivo* studies, 10 µM melatonin was used, along with high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM). Following the 30-minute drug pre-treatment, aortic contraction was induced through the cumulative administration of Phe (100 nM – 1 µM). Once maximum contraction was reach, aortic relaxation was induced through the cumulative administration of Ach (30 nM – 10 µM). *Ex vivo* treatment groups all contained a combined vehicle. Only endothelium-dependent *ex vivo* studies were conducted due to limited availability of rats, and well as due to the novel nature of these investigations. No L-NAME control experiments were included for the *ex vivo* studies due to limited equipment availability. For all *ex vivo* isometric tension studies, Phe-contraction results are expressed as an increase in gram tension from a resting tension of 1.5 g, while Ach-relaxation results are expressed as % relaxation of the maximum phenylephrine-pre-contraction. A p-value of < 0.05 was considered statistically significant.

5.1.1 Phe-contraction / Ach-relaxation

Cumulative contraction with Phe, showed that the ART treated aortas contracted (measured by increase in g tension from a stable point of 1.5 g) significantly less when compared to all other treatment groups (Two-way ANOVA Con: p=0.0070; Mel: p=0.0250; Mel + ART: p=0.0001 vs ART) (Fig 5.1 A).

The addition of Ach induced significantly less relaxation (measured by the % decrease in g tension from the point of maximal contraction) in the melatonin + ART group compared to the control group (Two-way ANOVA Con: p=0.0148 vs Mel+ART) (Fig 5.1 B).

No significant differences were seen in $E_{max}$ values between the different groups (See Table 5.1. for $E_{max}$ values). $E_{max}$ is the maximal response of the system.
Figure 5.1: Graphs indicating the effects of *ex vivo* melatonin and ART treatment on Phe induced contraction and Ach induced relaxation. A) Aortic ring contractions in response to cumulative concentrations of Phe; B) Aortic ring relaxation in response to cumulative concentrations of Ach (n = 5-6/group).
Table 5.1: Ex vivo $E_{\text{max}}$ values for the various treatment groups. Phe was used to induce contraction, and Ach was used to induce relaxation.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Melatonin</th>
<th>ART</th>
<th>Melatonin + ART</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phe (g) $E_{\text{max}}$ $\pm$ SEM</td>
<td>2.998 ± 0.1244</td>
<td>3.043 ± 0.1519</td>
<td>2.833 ± 0.1369</td>
<td>3.171 ± 0.1598</td>
</tr>
<tr>
<td>Ach (%) $E_{\text{max}}$ $\pm$ SEM</td>
<td>78.56 ± 4.682</td>
<td>67.47 ± 4.275</td>
<td>75.50 ± 4.685</td>
<td>61.01 ± 4.572</td>
</tr>
</tbody>
</table>

5.2 In Vivo Studies

The main aim of the in vivo studies was to determine the chronic effects of melatonin and ART administration on the vascular reactivity of male wistar rats that have been treated for 8 weeks with melatonin and/or ART. Additional aims included examining various protein activation and/or expression levels, as well as the antioxidant capacity exhibited in the aortic tissue of these same male Wistar rats.

For these in vivo studies, 80 male Wistar rats were randomly divided into four treatment groups: control, melatonin (10 mg/kg/day), ART (EFV: 51.6 mg/kg; FTC: 17.4 mg/kg; TDF: 25.8 mg/kg) and melatonin + ART. The rats received the respective treatments for 8 weeks, after which the animals were sacrificed, the aortas removed and the following data were obtained.

A p-value of < 0.05 was considered statistically significant.

5.2.1 Endothelium-dependent aortic ring investigations

As mentioned in section 3.3.3.2, two types of aortic ring investigations were conducted. The first set of aortic ring investigations conducted, were endothelium-dependent relaxation studies. For these studies, Phe was used to induce contraction and Ach was used to induce relaxation (as previously described above in section 5.1). As a negative control for the endothelium-dependent relaxation studies, some of the rings were pre-treated with the NOS inhibitor, L-NAME (100 µM) 15 minutes prior to the administration of cumulative amounts of Phe and Ach in order to confirm that the relaxation responses were indeed derived from endothelial NOS-NO release.
5.2.1.2 Phe-contraction / Ach-relaxation

Cumulative contraction with Phe, showed that the ART and melatonin + ART treatment groups contracted significantly more overall, when compared to the control group (Two-way ANOVA Con: p<0.0001 vs ART; Mel+ART) (Fig 5.2 A).

Ach induced relaxations were similar for all treatment groups (Fig 5.2 B).

No significant differences were seen in $E_{\text{max}}$ values between the different groups (See Table 5.2. for $E_{\text{max}}$ values).

Figure 5.2: Graphs indicating the effects of in vivo melatonin and ART treatment on Phe induced contraction and Ach induced relaxation. A) Aortic ring contractions in response to cumulative concentrations of Phe; B) Aortic ring relaxation in response to cumulative concentrations of Ach (n = 12-16/group).
Table 5.2: *In vivo* $E_{\text{max}}$ values for the various treatment groups. Phe was used to induce contraction, and Ach was used to induce relaxation.

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>Melatonin</th>
<th>ART</th>
<th>Melatonin + ART</th>
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<tbody>
<tr>
<td><strong>Phe (g)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$E_{\text{max}} \pm$ SEM</td>
<td>2.635 ± 0.07303</td>
<td>2.862 ± 0.07496</td>
<td>2.852 ± 0.07322</td>
<td>2.851 ± 0.07378</td>
</tr>
<tr>
<td><strong>Ach (%)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$E_{\text{max}} \pm$ SEM</td>
<td>79.80 ± 3.656</td>
<td>81.49 ± 3.662</td>
<td>84.34 ± 3.568</td>
<td>77.72 ± 3.878</td>
</tr>
</tbody>
</table>

5.2.1.3 L-NAME + Phe-contraction / Ach-relaxation

**Phe Contraction Studies**

Pre-treatment with 100 µM L-NAME caused all treatment groups (Two-way ANOVA $p<0.0001$ in all groups) to contract significantly more when compared to their respective L-NAME treated group (Fig 5.3 A). The L-NAME+control (3.714 ± 0.2148g) and L-NAME+melatonin (3.890 ± 0.1897g) groups experienced a ± 35% increase in contraction at the maximum Phe concentration compared to the control (2.761 ± 0.07774g) and melatonin (2.876 ± 0.08195g) groups, respectively (Fig 5.3 A). The L-NAME+ART (4.021 ± 0.2713g) group experienced a similar increase in maximum contraction of ± 36% when compared to the ART (2.953 ± 0.07830g) group. L-NAME+Mel+ART experienced a slightly smaller, but still noteworthy increase in maximum contraction of ± 29% when compared to the Mel+ART (2.950 ± 0.07860g) group (Fig 5.3 A).

However, when L-NAME pre-treated groups were compared with each other, there were no differences seen in Phe induced contraction (Fig 5.3 B), between any of the treatment groups.
Figure 5.3: The effects of L-NAME pre-administration on Phe-induced contraction. A) The differences in contraction of aortic rings between treatment groups incubated with or without L-NAME. B) The differences in contraction of aortic rings between treatment groups pre-treated with L-NAME (n = 4-5/group).
Ach Relaxation Studies

**Figure 5.4 A** shows that all treatment groups exposed to L-NAME pre-administration relaxed significantly less (Two-way ANOVA p<0.0001 in all groups) when compared to the applicable treatment groups not exposed to L-NAME pre-treatment. The L-NAME+control (0.7550 ± 0.7187%) and L-NAME+ART (1.191 ± 0.6933%) groups experienced a ± 99% decrease in relaxation capacity at the maximum Phe concentration compared to the control (79.80 ± 3.656%) and ART (84.34 ± 3.568%) groups, respectively (Fig 5.4 A). The L-NAME+melatonin (-1.479 ± 0.7575%) and L-NAME+Mel+ART (-1.245 ± 0.7238%) groups both continued to contract, even following the addition of the full cumulative dose of Ach (Fig 5.4 A). This meant the final tension (g) after the cumulative addition of Ach in these two groups, was higher when compared to the final tension seen after the accumulative addition of Phe. When the final mean tension values are compared (after the accumulative addition of both Phe and Ach), we see that L-NAME+melatonin had a high ending mean tension of 2.21g, while the melatonin group had a much lower ending mean tension of 0.25g. The same trend was seen in the L-NAME+melatonin+ART group, which had a final mean tension of 2.15g, compared to the much lower final mean tension of 0.31g in the melatonin+ART.

When Ach induced relaxation in L-NAME pre-treated groups were compared with each other, there were once again no differences seen (Fig 5.4 B), between any of the treatment groups. Some aortas continued to contract after the addition of Ach following pre-treatment with L-NAME, which consequently resulted in negative $E_{\text{max}}$ values.

No significant differences were seen in $E_{\text{max}}$ values between the different groups (See **Table 5.3** for $E_{\text{max}}$ values).
Figure 5.4: The effects of L-NAME pre-administration on Ach-induced relaxation. A) The differences in relaxation of aortic rings between treatment groups incubated with or without L-NAME. B) The differences in relaxation of aortic rings between treatment groups pre-treated with L-NAME (n = 4-5/group).
**Table 5.3:** *In vivo* $E_{\text{max}}$ values for L-NAME pre-treated treatment groups. Phe was used to induce contraction, and Ach was used to induce relaxation.

<table>
<thead>
<tr>
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<th>Melatonin + ART</th>
</tr>
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<tbody>
<tr>
<td><strong>Phe (g) $E_{\text{max}} \pm \text{SEM}$</strong></td>
<td>3.714 ± 0.2148</td>
<td>3.890 ± 0.1897</td>
<td>4.021 ± 0.2713</td>
<td>3.793 ± 0.1767</td>
</tr>
<tr>
<td><strong>Ach (%) $E_{\text{max}} \pm \text{SEM}$</strong></td>
<td>0.7550 ± 0.7187</td>
<td>-1.479 ± 0.7575</td>
<td>1.191 ± 0.6933</td>
<td>-1.245 ± 0.7238</td>
</tr>
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</table>

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**5.2.2 Endothelium-independent aortic ring investigations**

The second set of aortic ring investigations conducted, were endothelium-independent relaxation studies. For these studies, Phe was once again used to induce contraction, however, SNP - an exogenous NO donor - was used to induce relaxation.

**5.2.2.1 Phe-contraction / SNP-relaxation**

Cumulative contraction with Phe, showed that the ART and melatonin + ART treatment groups exhibited significant pro-contractile properties compared to the control group (Two-way ANOVA Con: p<0.0001 vs ART; Mel+ART) (Fig 5.5 A).

SNP induced relaxations were similar for all treatment groups (Fig 5.5 B).

No significant differences were seen in $E_{\text{max}}$ values between the different groups (See Table 5.4. for Emax values).
Figure 5.5: Graphs indicating the effects of in vivo melatonin and ART treatment on Phe induced contraction and SNP induced relaxation. A) Aortic ring contractions in response to cumulative concentrations of Phe; B) Aortic ring relaxation in response to cumulative concentrations of SNP (n = 5/group).
Table 5.4: In vivo $E_{\text{max}}$ values for the various treatment groups. Phe was used to induce contraction, and SNP was used to induce relaxation.

<table>
<thead>
<tr>
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<td>3.793 $\pm$ 0.1767</td>
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<tr>
<td>Ach (%)</td>
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<td></td>
</tr>
<tr>
<td>$E_{\text{max}}$ $\pm$ SEM</td>
<td>110.3 $\pm$ 4.639</td>
<td>113.2 $\pm$ 3.860</td>
<td>108.1 $\pm$ 4.153</td>
<td>108.0 $\pm$ 4.130</td>
</tr>
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</table>

5.2.3 Western Blot Analyses

Western blot analyses were performed on snap-frozen aortic tissue of control, melatonin, ART and melatonin + ART treated animals in order to determine whether the functional data of the vascular reactivity investigations could be explained by changes in intracellular signal transduction pathways.

All blots are calculated and expressed as a ratio of the corresponding loading controls. Controls are expressed as 1.

5.2.3.1 NO & RNS Signalling

**eNOS**

eNOS phosphorylation was significantly lower in the melatonin + ART (0.3292 $\pm$ 0.04093) treatment group compared to the control (1.000 $\pm$ 0.2814). No significant differences were seen in the phosphorylation levels of the melatonin (0.5541 $\pm$ 0.06424) or ART (0.5251 $\pm$ 0.1112) treatment groups (Fig 5.6 A).

No significant differences were seen in total eNOS expression between any of the treatment groups (Mel: 0.6192 $\pm$ 0.09478; ART: 0.5239 $\pm$ 0.07665; Mel + ART: 0.3522 $\pm$ 0.07480) when compared to the control (1.000 $\pm$ 0.3073) (Fig 5.6 B).

When eNOS P/T ratios were calculated, no significant differences were seen in total eNOS expression between any of the treatment groups (Con: 1.000 $\pm$ 0.04927; Mel: 1.081 $\pm$ 0.06813; ART 1.156 $\pm$ 0.07625; Mel+ART: 1.187 $\pm$ 0.1149) (Fig 5.6 C).
Figure 5.6: Changes in the eNOS phosphorylation and expression (Ser 1177) in aortic tissue from rats treated with melatonin and ART, in vivo. A) Analysed results and western blot for phospho-eNOS (Ser 1177); B) Analysed results and western blot for total-eNOS; C) Phosphorylated over total (P/T) ratio of eNOS (Ser 1177) (n=5/group).
iNOS

No significant differences were seen in total iNOS expression between any of the treatment groups (Fig 5.7)

**Figure 5.7:** Changes in iNOS expression in aortic tissue from rats treated with melatonin and ART, *in vivo*. Shown are the analysed results and western blot for iNOS (n=5/group).
Nitrotyrosine

No significant differences were seen in nitrotyrosine levels between any of the treatment groups (Fig 5.8)

Figure 5.8: Changes in nitrotyrosine expression in aortic tissue from rats treated with melatonin and ART, *in vivo*. Shown are the analysed results and western blot for nitrotyrosine (n=5/ group).
5.2.3.2 ROS Signalling

p22 PHOX

No significant differences were seen in total p22 PHOX expression between any of the treatment groups (Fig 5.9).

Figure 5.9: Changes in the p22 PHOX expression in aortic tissue from rats treated with melatonin and ART, *in vivo*. Shown are the analysed results and western blot for p22 PHOX (n=5/group).
5.2.3.3 Cell Viability Signalling

Cleaved Caspase-3

The ART (0.2735 ± 0.08317) and melatonin + ART (0.4757 ± 0.08704) combination treatment groups showed significantly decreased cleaved Caspase-3 expression compared to the control (1.000 ± 0.1945). The ART treatment group also showed significantly decreased cleaved Caspase-3 expression when compared to the melatonin treatment group (0.6309 ± 0.09441) (Fig 5.10).

![Cleaved Caspase-3 expression graph]

**Figure 5.10:** Changes in cleaved Caspase-3 expression in aortic tissue from rats treated with melatonin and ART, in vivo. Shown are the analysed results and western blot for cleaved Caspase-3 (n=5/group).
5.2.4 Antioxidant Capacity Studies

Antioxidant capacity studies were performed on snap-frozen aortic tissue of control, melatonin, ART and melatonin + ART treated animals.

Antioxidant capacity is expressed in trolox equivalents, which is a vitamin-E analogue that is used as the standard measure of antioxidant capacity.

5.2.4.1 ORAC Assay

No significant differences were seen between melatonin (0.0564 ± 0.01664), ART (0.0532 ± 0.01131) and melatonin + ART (0.07225 ± 0.006700) treatment groups when compared to the control (0.0430 ± 0.005899) (Fig 5.11).

![Graph showing antioxidant capacity of aortic tissue from rats treated with melatonin and ART, in vivo (n = 5/group).](Figure%205.11%3A%20Antioxidant%20capacity%20of%20aortic%20tissue%20from%20rats%20treated%20with%20melatonin%20and%20ART,%20in%20vivo%20(n%20=%205/group).)
6 – Discussion: *In vitro* studies

Chapter 6 debates the findings of the results obtained from the *in vitro* study. The data discussed in this chapter can be found in Chapter 4, while the materials and methods can be found in Chapter 2.

Throughout the *in vitro* studies, we set out to examine three important variables: NO and RNS levels, ROS levels and cell viability. All three of these factors play important roles in the maintenance of endothelial health and vascular function (Mudau et al. 2012; K. Park & Park, 2015; Versari et al., 2009).

NO is viewed as an endothelial cell survival factor (Dimmeler & Zeiher, 1999), and a reduction in NO bioavailability is considered as a marker for ED (Muniyappa & Sowers, 2013). However, besides its important beneficial activities, in pathophysiological conditions excessive NO production may contribute to oxidative stress and an associated inflammatory state with eventual tissue damage (Pacher et al., 2007). Together, excessive NO or RNS, as well as ROS, can cause a decrease in cell viability and eventually lead to cell death (Circu & Aw, 2010; Fransen et al., 2012; Marín-García, 2016; Murphy, 1999). For cell viability and cell death, we examined both necrosis and apoptosis. Necrosis refers to the pathological or accidental mode of cell death involving irreversible swelling of the cytoplasm and distortion of organelles including mitochondria followed by rupture of the cell membrane and inflammation that damage the cells and its surrounding tissues (Fadeel & Orrenius, 2005). Apoptosis is a sequential process where the dying cell undergoes nuclear and cytoplasmic condensation with blebbing of the plasma membrane leading to formation of apoptotic bodies, which are recognised and removed, via phagocytosis by macrophages without damaging the surrounding tissues (Fadeel & Orrenius, 2005).

6.1 Melatonin Dose-Response

The antioxidant activity of melatonin has been reported at both physiological and pharmacological concentrations (Galano et al., 2011; Reiter et al., 2005), although pharmacological dosages of exogenously administered melatonin typically provide greater protection from large quantities of free radicals (Tan et al., 2007). In literature, *in vitro* melatonin supplementation studies conducted on various types of ECs have used a wide
range of melatonin concentrations. Some studies have used dosages as low as 0.1 nM (Tamura, Silva, & Markus, 2006), while others have used concentrations as high as 50 µM (Pogan, Bissonnette, Parent, & Sauve, 2002). However, most studies used a concentration range between 1 nM and 10 µM (Silva et al., 2007; Tamura, Cecon, Monteiro, Silva, & Markus, 2009; Urata et al., 1999), and therefore in this study, we decided to use 1 nM; 1 µM and 10 µM melatonin in the dose-response studies.

None of the three melatonin dosages tested, elicited any changes in NO (Fig 4.1 & 4.4) or RNS production (Fig 4.2 & 4.5), regardless of the treatment period (1h or 24h). This is contradictory to most in vitro findings in literature, where these concentrations of melatonin have been shown to decrease NO and RNS production (Jumnongprakhon, Govitrapong, Tocharus, & Tocharus, 2016; Reiter et al., 2001; Silva et al., 2007; Tamura et al., 2006; Tamura et al., 2009). However, the abovementioned studies were conducted using rat brain microvascular endothelial cells as well as rat microvascular cremaster muscle endothelial cells. Although these are also endothelial cell lines, the function of endothelial cells are specialised depending on their location (Gordon et al., 1991), making a direct comparison difficult. On the other hand, and in line with this investigation, a study by Tamura et al. (2006) found that the treatment of rat microvascular endothelial cells with 0.1 nM and 1 nM melatonin had no effect on NO production. However, the observed result in the aforementioned study is most likely due to the very short melatonin incubation period of 1 minute. No studies have been conducted with the specific ECs examined in this study, namely AECs. Therefore further studies would be needed to determine whether the lack of changes in NO and ROS levels observed in AECs, as shown in these results, is a standard response for an AEC, or if these findings are the result of the specific technique used. Various techniques are available to measure NO and RNS production, however, the low levels of NO and RNS produced by cultured vascular ECs in vitro exposes the sensitivity limits of many of these assays (Kleinhizen et al., 2003). In this case, a potential more sensitive alternative to fluorometric detection of NO and RNS may be to use a chemiluminescence technique, which has shown to detect these molecules at concentrations as low as 1 p.m., compared to the 2-5 nM detection capacity of the employed fluorometric method (Kleinhizen et al., 2003).

Conversely, the cell viability studies in the present study showed that 1 nM melatonin was the only concentration to decrease necrosis after a 24-hour treatment period (Fig 4.3),
which was in line with findings from other studies (Harms et al., 2000; Song et al., 2014). It was however expected that all melatonin concentrations would decrease necrosis, since the examined melatonin concentration range (1 nM; 1 µM and 10 µM) has been shown to increase cell viability over and incubation period of 24h, in multiple other studies (Celik & Maziroglu, 2012; Kim et al., 2011; Osseni et al., 2000). The lack of melatonin activity in the 1 µM and 10 µM treatment groups after 24 hours of treatment was therefore very surprising. In contrast, a study conducted on a human liver cell line (HepG2), showed that melatonin treatment at concentrations of 1-10,000 µM over a longer treatment period (24–96 h) decreased cell viability (Osseni et al., 2000). This finding is line with another study conducted in our laboratory, which found that cardiac microvascular endothelial cells (CMECs) showed decreased cell viability after 6-24 h melatonin treatment (1 nM; 1 µM and 10 µM melatonin) (Nduhirabandi, 2014). In the current study, AECs showed no significant loss in cell viability after treatment with higher concentrations of melatonin over 24h (Fig 4.3), which may indicate that AECs are more robust than other previously investigated cell lines. The robustness of this cell line may also be a possible explanation for the lack of significant changes seen in the NO and RNS investigations. A potential solution may be to apply a harmful but not lethal stimulus to the cells before treatment, in order to elicit a protective response by melatonin in otherwise healthy cells.

After the 1-hour treatment period, all concentrations of melatonin showed a significant decrease in necrosis compared to the control, with the addition of 1 nM melatonin decreasing necrosis significantly more compared to 10 µM melatonin (Fig 4.6). The study by Nduhirabandi (2014) mentioned in the paragraph above, also showed that a melatonin treatment (1 nM; 1 µM and 10 µM) period of 1 hour, was the only treatment period where melatonin increased cell viability. The current study’s finding that melatonin increases cell viability most effectively at low concentrations over a short treatment period may be explained by a finding that a higher melatonin concentration (10 µM) induced GSH depletion, that was associated with ROS over-production after only 15 minutes of incubation (Osseni et al., 2000).

Overall, these melatonin dose-response findings led us to conclude that a concentration of 1 nM melatonin would yield the best results in the future combination studies, since it was the only dose where significant differences were observed for the 24 hour treatment period, and the most effective dose over the 1 hour treatment period.
6.2 ART Dose-Response

Despite confounding reports, many trials indicate that ART leads to ED (Skowyra et al., 2012). Cardiovascular complications are considered a new challenge for HIV patients receiving ART with numerous studies on the effects of ART on the vasculature indicating that ART may impair the function of the vascular endothelium (Blanco et al., 2006; Shankar et al., 2005; Stein et al., 2001). ART has been shown to affect NO levels; either through the excessive production of NO (Torre et al., 2015) or through reduced NO production (Shankar et al., 2005), due to reduced expression of eNOS (Fu et al., 2005). Increases in ROS and RNS production (Skowyra et al., 2012) have also been demonstrated in ART, and together, these factors can lead to decreased cell viability.

In this study, high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) significantly increased NO production compared to both the control and the other two ART concentrations (Fig 4.7). These finding of increased NO production during treatment with ART were surprising. Although it has been reported in literature that ART has the ability to increase excessive production of NO (Torre et al., 2015), the study conducted by was conducted in humans, making it difficult to compare these results with a cell-based study. NO is a cell survival factor (Dimmeler & Zeiher, 1999), and its controlled production plays many important roles in the immune and vascular system (Cines et al., 1998). In the case of cell injury and the release of inflammatory mediators, iNOS is responsible for producing characteristically high concentrations of NO in shorts bursts in order to protect the cells (Matthys & Bult, 1997; Michel & Feron, 1997). Under certain circumstances, this high concentration of NO produced by iNOS can become detrimental, and excessive NO may damage the vascular wall (Matthys & Bult, 1997) or cause cell death through oxidative stress (Murphy, 1999). In the current study, the increase in NO observed between the control group and the highest ART concentration treatment group was as modest as ±13% (Fig 4.7). A study by Speyer et al., (2003) showed that iNOS induced NO production was 20-fold above that found in non-stimulated endothelial cells, where iNOS was inactive. Therefore, it is more likely that the modest increase in NO levels seen in the current study are more likely to be associated with a protective effect on the endothelium, rather than a damaging effect.

Clinical and experimental models have shown that ART-induced ED appears to be mediated by reduced NO production (Shankar et al., 2005). Most these clinical and experimental models make use of protease inhibitors, as opposed to NRTIs and NNRTIs, which comprise first line FDC ART used in the current study. The fact that this study
observed increased NO production after treatment with NRTI and NNRTI based ART may imply that NRTI and NNRTI based therapies may be a safer treatment option with regards to endothelial health and maintaining NO homeostasis. Alternatively, the result of this study may be due to the relatively short incubation time of 24-hours. A longer treatment period is more applicable when treating with ART, since ART-associated ED is usually only observed after many years of treatment (Choi et al., 2011; Islam et al., 2012). Further studies would need to be conducted to determine if NO production would still increase after a longer treatment period.

When RNS production was examined, high concentration ART was the only concentration to significantly increase RNS production (Fig 4.8), with the increase being a slight as ±9% higher than the control group. It is not surprising that there were no significant differences seen in the two lower concentration ART groups, as well as that the high concentration ART group only resulted in a minor increase in RNS production, since no major increase in NO production was observed either. These two finding are related, since excessive levels of NO are associated with increased ONOO\(^{-}\) levels (Forstermann & Munzel, 2006), which in turn is associated with the generation of nitrotyrosine (Halliwell, 1997). Therefore, in the absence of excessive NO production, excessive production of RNS is also unlikely. One study examined the effects of ART in a hCMEC/D3 cell line (in vitro model for blood-brain barrier endothelial cells) and found ROS to be highly elevated post ART treatment (Manda, Banerjee, Banks, & Ercal, 2011). However, as is the case with most in vitro studies done on the effects of ART, this study only examined ROS production and not RNS production, as well as used a longer treatment period of 72 hours. Multiple studies have demonstrated increased ART-induced ROS production (Apostolova et al., 2010; Skowyra et al., 2012; Weiß et al., 2016), but no studies have looked at the effects of ART treatment on RNS (specifically peroxynitrite) in vitro. It is therefore important to further investigate the role of RNS in ART-induced ED, by testing different classes of ARV drugs as well as using different techniques to detect specific types of RNS to verify a potential source of RNS.

For the cell viability studies, high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) increased necrosis in AECs, when compared to the control (Fig 4.9). High concentration ART also showed significantly increased necrosis compared to the other two ART concentrations (Fig 4.9). This results correspond with the RNS production results (Fig 4.8), where we saw that the high concentration ART group has the largest increase in RNS.
production, which could potentially, in part, be responsible for the decreased cell viability observed, since increases in RNS are associated with cell death (Murphy, 1999).

These results are comparable to a study that demonstrated increased necrosis in human umbilical vein endothelial cells (HUVECs) following treatment with EFV (Weiß et al., 2016), one of the components of first line FDC ART. The above-mentioned study treated cells with 30 – 50 µM EFV. The current study has demonstrated a similar increase in necrosis with a much lower concentration of 12 µM EFV. Another study by Zhong et al. (2002) showed that protease inhibitor ritonavir directly caused endothelial cell death, mainly through necrosis pathways in human dermal microvascular endothelial cells (HMECs). The only other study conducted on AECs that examined the effect of first line FDC ART treatment on cell viability was conducted in our laboratory. The mentioned study showed that first line FDC ART has no effect on cell viability, at any tested concentration, compared to the untreated control group (Charania, 2017). This finding is inconsistent with the findings of this study, but this discrepancy can most likely be attributed to the lower ART concentration ranges used in the other study (maximum ART concentrations – EFV: 11.2 nM; FTC: 2.6 µM; TDF: 1 µM).

From our findings, we determined that high concentration ART would be the most effective dosage for the subsequent experiments, since it led to the greatest changes in NO- and RNS production, as well as necrosis.

### 6.3 Effects of melatonin and ART treatment on NO production, RNS production and cell viability in AECs

As mentioned above in section 6.1 and 6.2, from the dose-response studies, it was determined that 1 nM melatonin along with high concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) would be used in the combination studies.

In the NO- and RNS production combination studies, no significant differences were observed between any of the treatment groups (Fig 4.10 and 4.11). This finding was not very surprising with regards to the melatonin groups, since in the dose response studies, melatonin treatment had no effect on NO- and RNS-production. But for the ART treatment groups, we did expect the results from the dose-response studies to be mirrored in the combination treatment investigations. One explanation for these contrasting results may be that new DAF-2/DA and DHR-123 batches had to be purchased and used for the combination experiments. However, it is unlikely that this change should affect the results, since the probes were bought from the same supplier. It is therefore difficult to explain the
discrepancy, as the experimental conditions (other than the different fluorescent probe batches), were identical to the dose-response studies.

Results from the cell viability studies were more in line with the dose-response findings. We found that 1 nM melatonin significantly decreased necrosis compared to the control, while high concentration ART significantly increased necrosis compared to both the control and the 1 nM melatonin treatment group (Fig 4.12). When the two drugs were combined, we saw a significant decrease in necrosis for the combination group compared to the group treated with ART alone (Fig 4.12). In this way, we demonstrated that melatonin was able to ameliorate the harmful effects of ART. This is a novel finding, since to the best of our knowledge, there are no studies that have demonstrated this in AECs using first line FDC ART and melatonin.

These observations may be due to melatonin’s antioxidant capabilities, which would allow the cells to remain viable in situations of stress where toxins like ART may lead to a loss in cell membrane viability. However, more specific investigations into the underlying mechanisms are necessary to determine through what means melatonin is able to induce these protective changes.

6.4 Western Blot Analyses

6.4.1 NO & RNS Signalling

It is clear that NO and NOS play key roles in endothelial function, not only in the context of inducing vasodilatation, but also through a plethora of other important regulatory actions (Cines et al., 1998). It is essential for the healthy endothelium that both NOS (as the most important source of vascular NO), and NO are present in desirable quantities, since an imbalance in NO and NOS levels can have dire consequences for endothelial health.

6.4.1.1 eNOS

eNOS is the most abundant isoform of NOS and is responsible for the NO produced in the vasculature, and therefore vessel dilation (Sandoo et al., 2010).

Since no changes in NO production were observed with the fluorescent probe assays, it was expected that eNOS activation (phosphorylation) (Fig 4.13 A) and expression (total protein) levels (Fig 4.13 B) would also remain unchanged. When the phosphorylated eNOS was expressed as a ratio of total, the ratio of all treatment groups decreased compared to the control group, while the melatonin + ART group also decreased significantly compared to the melatonin group (Fig 4.13 C). This ratio was seen to be as a
result of decreased phosphorylated eNOS levels as opposed to increased total eNOS levels. It is well reported that melatonin has the ability to decrease NO levels through its antioxidative capacity (Jumnongprakhon et al., 2016; Reiter et al., 2001; Silva et al., 2007; Tamura et al., 2009). These convincing Western blot results suggest that melatonin treatment could decrease NO levels through decreased eNOS activation. It was interesting, however, that the groups treated with ART also showed decreased eNOS activation, whilst the dose-response studies showed that ART increased NO production. This may imply that ART-induced NO production may be independent of the endothelium and eNOS, although further studies would need to be done for further elucidation on this point.

6.4.1.2 iNOS

While eNOS is associated with the maintenance of basal levels of NO, iNOS has traditionally been associated with high output NO-release in response to inflammatory mediators (Buchwalow et al., 2001; Michel & Feron, 1997) and in many cases is associated with cytotoxic peroxynitrite formation (Strijdom et al., 2009).

iNOS was not detected in the AECs (Fig 4.14). This may imply that iNOS might not have been expressed or that iNOS levels were so low that the antibody was not able to detect iNOS in the cells. Both eNOS and iNOS can therefore be excluded as sources of the increased NO production observed. Therefore this increase in NO production seen in the dose-response studies still has an unknown source that needs investigating. Potential sources of NOS-independent NO include S-nitrosothiols, as well as nitrite and nitrate (Chen, Pittman, & Popel, 2008). Under certain conditions, nitrite and nitrate act as a reservoir, where different enzymes (hemoglobin, myoglobin, xanthine oxidoreductase, mitochondrial cytochrome oxidase, aldehyde dehydrogenase 2, cytochrome P450 reductase and cytochrome P450) can catalyze the reduction of nitrite or nitrate to generate NO (Zhao, Vanhoutte, & Leung, 2015). Various methods exist to measure NOS-independent NO formation (Zweier, Samouilov, & Kuppusamy, 1999), which could be examined in future studies.

6.4.1.3 Nitrotyrosine

Nitrotyrosine is generated when peroxynitrite reacts with a tyrosine or tyrosine containing protein and serves as a marker for nitrosative stress and peroxynitrite damage (Halliwell, 1997). Peroxynitrite is generated as a result of the chemical reaction between NO and superoxide (Forstermann & Munzel, 2006).
Nitrotyrosine levels in the ART treatment group were significantly higher when compared to the control group (Fig 4.15). This finding corresponds to the current literature, which suggests that ART leads to the excess production of ROS, particularly superoxide (Sutliff et al., 2002). Superoxide, along with increased levels of NO production leads to the production of peroxynitrite and eventually nitrotyrosine. The treatment group containing both melatonin and ART showed significantly lower levels of nitrotyrosine compared to the ART group (Fig 4.15). These results may be indicative that melatonin has the ability to decrease the nitrosative stress caused by ART. This protection by melatonin is most likely due to its antioxidative capacities and its ability to scavenge free radicals like superoxide and excess NO (Hardeland, 2005; Poeggeler et al., 2002), thereby decreasing the chance that peroxynitrite and nitrotyrosine will form (Fig 6.1). These results are similar to those seen in the plate reader studies, where ART significantly increased DHR-123 fluorescence compared to the control. Together, these results may suggest that ART induces ED through excess production of RNS like peroxynitrite. Most the literature covers the effects of ART treatment on ROS production. This study establishes that RNS may play and equally important role in the multifactor mechanisms by which ART induces ED. The plate reader studies did not however show similar results with regards to the alleviative effect of melatonin on RNS levels. This may be due to the fact that the nitrotyrosine antibody is a more specific and sensitive probe compared to DHR-123. Further protein and antioxidant capacity studies would be needed to confirm melatonin’s specific role in ameliorating nitrosative stress.

Figure 6.1: Substrates involved in nitrotyrosine production (Everson, 2016).

6.4.2 ROS Signalling

ART has been shown to result in increased ROS-production from various sources (Apostolova et al., 2010; Skowyra et al., 2012; Weiß et al., 2016). Superoxide is of
particular interest since it is essentially a precursor of other harmful radicals and molecules, such as hydrogen peroxide, peroxynitrite, the hydroxyl radical and nitrite.

6.4.2.1 P22 PHOX

p22 PHOX is one of multiple subunits of NADPH oxidase, which is responsible for superoxide production. p22 PHOX is therefore regarded as a marker of NADPH-oxidase dependent superoxide production (San Jose, Fortuno, Beloqui, Diez, & Zalba, 2008).

All treatment groups showed lower levels of p22 PHOX expression compared to the control group (Fig 4.16). Both the ART and melatonin + ART groups also showed significantly lower p22 PHOX expression compared to the melatonin group, potentially indicating that ART may be the factor lowering the p22 PHOX expression levels. These results contradict the findings of the DHR-123 investigations where ART increased RNS (peroxynitrite) production, as well as opposes many previous reports in literature. This inconsistent finding may be due to the fact p22 PHOX is specific to NADPH-oxidase dependent superoxide, while DHR-123 is reported to be more sensitive to peroxynitrite, although some studies have also shown it to be sensitive to mitochondrial ROS (Chan & Miskimins, 2012; Ischiropoulos et al., 1999; Navarro-Antolin et al., 2001). NADPH-oxidase has been implicated as the main source of excess ROS production in the vasculature (Hamilton et al., 2002), however, the main source of ART induced superoxide production is thought to be of mitochondrial origin, likely resulting from leakage of electrons from a dysfunctional mitochondrial electron transport chain (Jiang et al., 2007). The results from the DHR-123 study substantiate this, since the probe is more sensitive to mitochondrial ROS and RNS (peroxynitrite), and showed a marked increase in ART treated groups, while the p22 PHOX did not. A marker with greater specificity for mitochondrial superoxide would need to be evaluated. Superoxide dismutase 2 (SOD-2) is involved in the scavenging of superoxide that leaks from the mitochondrial electron transport chain (Zelko, Mariani, & Folz, 2002), and therefore could be an alternative target for further investigation.

6.4.3 Cell Viability Signalling

Previous in vitro investigations have examined cell viability through necrosis. However, apoptosis is also an important form of cell death which utilises a different protein transduction pathway to necrosis, and may also be involved in ART induced cell damage.

6.4.3.1 Cleaved Caspase-3
Cleaved caspase-3 is a down-stream target of caspase-8 and indicative of apoptosis (Kuribayashi et al., 2006).

The ART and melatonin + ART treatment groups both showed decreased cleaved caspase-3 expression levels compared to both the control and the melatonin treatment groups (Fig 4.17). In terms of cell viability, these results contradict the findings from the PI-based studies where we saw ART increased necrosis, although apoptosis and necrosis do not necessarily occur simultaneously and therefore cannot be directly compared. These confounding results correlate with evidence from literature, which demonstrates that ART directly induces mitochondrial dysfunction and ROS production that promotes ED without culminating in apoptosis (Jiang et al., 2007).

Another explanation for decreased cleaved caspase-3 expression in groups containing ART may involve ART-induced NO production, which was observed in the dose-response studies. NO has previously been shown to inhibit apoptosis by S-nitrosylation of caspase-3 (Rossig et al., 1999), leading to its inactivation.

ART has been shown to be anti-apoptotic in specific cell types (Badley, 2005; Dieye et al., 2000), although the exact molecular target responsible for this anti-apoptotic effect remains to be defined in vitro and in vivo (Badley, 2005). The same molecular mechanisms could be at work here, but further protein investigations are needed.

### 6.5 Antioxidant Capacity Analyses

The oxygen radical absorbance capacity (ORAC) assay was used to assess the antioxidant capacity of the AECs treated with melatonin and ART. Unfortunately, no significant differences were seen between any treatment groups (Fig 4.18). It has been suggested that antioxidant activity should not be concluded based on a single antioxidant test model (Alam, Bristi, & Rafiquzzaman, 2013). Therefore, it might be prudent to rather conduct multiple more specialised and specific assays, such as hydrogen peroxide-; nitric oxide- and peroxynitrite radical scavenging assays, as well as thiobarbituric acid reactive substances (TBARS) assays which indicate lipid peroxidation.

### 6.6 Concluding Remarks

The first main aim for this in vitro study was to determine the optimum concentrations of both melatonin and first line FDC ART used to treat AECs at which the greatest differences could be seen when compared to untreated AECs, with respect to NO- and ROS production, as well as cell viability. It was established that 1 nM melatonin and high
concentration ART (EFV: 12µM; FTC: 10µM; TDF: 1µM) showed the greatest differences with regards to examined variables. Unfortunately, the melatonin dose could only be based on results from cell viability studies since no significant differences were seen in the NO- and ROS production studies.

The second main aim was to analyse any potential synergistic/antagonistic interactions between melatonin and ART in AECs treated with the above-determined concentrations, with respect to NO- and ROS production, as well as cell viability. For this aim, the NO- and ROS production studies did not show the same trends previously established in the dose-response studies. However, the changes observed in the dose-response studies were very modest, and would have perhaps been mirrored in the combination studies after an increase in the sample size. Cell viability studies were nevertheless followed the same trends observed in the dose-response studies and showed that melatonin alleviated necrotic damages induced by ART.

Signalling protein investigations established that the observed ART-induced increase in NO production might be independent of the endothelium and eNOS, as well as iNOS, while melatonin was shown to have the capacity to ameliorate nitrosative damage. p22 PHOX signalling eliminated NADPH-oxidase as a potential source of ROS in the ART-induced increase in ROS production and cleaved caspse-3 investigations showed that ART may by pro-necrotic, but it is most likely anti-apoptotic. Antioxidant capacity studies showed no significant changes between treatment groups. Overall, melatonin was shown to be effective in reducing cell necrosis and nitrosative damage, while ART may increase NO- and ROS production, and reduce apoptosis, but the mechanisms behind these changes still need further investigation.
7 - Discussion: Ex vivo and in vivo studies

Chapter 7 debates the findings of the results obtained from the ex vivo and in vivo studies. The data discussed in this chapter can be found in Chapter 5, while the materials and methods can be found in Chapter 3.

7.1 Ex Vivo Studies

The main aim of the ex vivo studies was to determine the acute effects of melatonin and ART administration on the vascular reactivity of untreated, control male Wistar rat aortas. For these studies, Phe was used to induce contraction, while Ach was used to induce relaxation. The relevant treatment was added directly to an ex vivo organ bath for 30-minutes before the addition of cumulative concentrations of Ach and Phe.

Cumulative contraction with Phe, showed that the ART treated aortas contracted significantly less, when compared to all other treatment groups (Fig 5.1 A). This finding may be the consequence of an unexpected phenomenon that occurred when ART was administered to the ex vivo organ bath (Fig 7.1). Roughly 1 minute after the addition of ART to the organ bath, the aortic ring started to spontaneously contract. On average it took about 5 minutes to reach maximum spontaneous contraction, before the ring began to relax again, eventually relaxing all the way back to a resting tension of 1.5 g before the end of the 30-minute ex vivo drug incubation period (Fig 7.1). When the vehicle or melatonin was added to the organ bath, no spontaneous contraction occurred.
It is possible that this pre-contraction caused by ART may have pre-conditioned the aortic ring to contraction, thereby causing it to contract less intensely when exposed to cumulative concentrations of Phe. Melatonin had the ability to attenuate the pre-conditioning effect of ART, since the melatonin + ART group contracted significantly more than the ART group during the subsequent cumulative Phe protocol, despite also being exposed to a spontaneous pre-contraction following *ex vivo* drug treatment. It is difficult to speculate how melatonin achieves this, since the mechanism of the ART-induced spontaneous pre-contraction is not known either. The spontaneous pre-contraction may be a result of iNOS activation following a bombardment of ROS generated by the *ex vivo* addition of ART. iNOS activation would lead to an excessive release of NO, which in combination with ROS, would be rapidly converted to peroxynitrite and ultimately lead to vascular contraction. Acute ART treatment might also directly stimulate pro-contractile substances such as endothelin-1 (ET-1), thromboxane A (TXA2) and angiotensin II (Cines et al., 1998; Mudau et al., 2012). This may however be slightly more difficult to examine, although an ELISA assay may be able to detect these substances within the supernatant. Alternatively, the spontaneous contraction may just be a direct drug-induced effect due to an unknown toxicity of one of the components of the FDC ART. Individual components of the ART (EFV; FTC and TDF) could be examined separately to clarify this point. Finally, the actuate treatment of ART might stimulate α-adrenergic receptors to release calcium,

![Figure 7.1](image.png)

**Figure 7.1:** The spontaneous contraction of *ex vivo* aortic rings after the addition of A) ART and B) ART and melatonin.
thereby inducing the rapid contraction. This rapid stimulation and excessive release of calcium may exhaust calcium supplies, which would explain the decreased contraction experience on administration of Phe.

However, this is all speculative and further investigations are needed to determine the source of this interesting and unexpected phenomenon, as well as the means by which melatonin attenuated the pre-conditioning effect of ART.

Further protein investigations should be conducted, as well as studies conducted with Ca\(^{2+}\) channel blockers, to further elucidate the mechanisms at play. The sample sizes should also be expanded to shed more light on these findings.

**7.2 In Vivo Studies**

The main aim of the *in vivo* studies was to determine the chronic effects of melatonin and ART administration on the vascular reactivity of male Wistar rats that have been treated for 8 weeks with melatonin and/or ART. Additional aims included examining various signalling protein activation and/or expression levels, as well as the antioxidant capacity exhibited in the aortic tissue of these same male Wistar rats.

For *in vivo* experiments, two types of aortic ring investigations were conducted. For the first set of experiments we conducted endothelium-dependent relaxation studies, where Phe was used to induce contraction and Ach was used to induce relaxation, and rings were pre-treated with the NOS inhibitor, L-NAME as a control. For the second set of experiments we conducted endothelium-independent relaxation studies, where Phe was once again used to induce contraction, however, for these studies SNP - an exogenous NO donor - was used to induce relaxation.

**7.2.1 Endothelium-dependent Aortic Ring Investigations**

Cumulative contraction with Phe, showed that the ART and melatonin + ART treatment groups contracted significantly more when compared to the control group (Fig 5.2 A). These results may implicate ART as the factor leading to increased contractility. Two studies by Jiang *et al.* (2006 and 2010) found that ART treatment had no effect on Phe-induced contraction. Both these studies used NRTI, azidothymidine (AZT) and the protease inhibitor, indinavir. Tests were also done on mouse aortas rather than rat aortas. Since a different animal model was used, as well as different ART, these two studies cannot be directly compared with the current study, and therefore the contradictory results are not unexpected. However, FTC, one of the components of first line FDC ART, is also a
nucleoside reverse transcriptase inhibitor, like AZT. Perhaps a conclusion that can be
drawn from the above-mentioned two studies is that the NRTI (FTC) component of the
combination ART is not responsible for the pro-contractile effects seen.

Studies examining the effects of ART treatment on aortic contractility are scarce and it is
therefore difficult to determine what might be the cause of ART-induced pro-contractility.
Further protein examination would be needed to confirm this mechanism. A potential
mechanism for this pro-contractile effect may be ART-induced reduced eNOS expression,
and therefore reduced NO release from the endothelium. Since we saw reduced eNOS
expression in the ART treated groups in vitro, the same might occur in vivo.

The fact that the ART groups experienced increased contractility in vivo is an expected
finding, since ART in vitro is thought to lead to impaired function of the vascular
endothelium (Blanco et al., 2006; Shankar et al., 2005; Stein et al., 2001). This is however
a contradictory result when compared to the observations made in the ex vivo studies,
where we saw ART treated groups showed anti-contractile properties. The only
explanation fathomable for these opposing results is that the duration of exposure to ART
plays a significant role in the resultant contractility effect. In other words, the longer the
endothelium is exposed to ART for, the greater the risk of developing ED. This theory has
been supported by many studies, with the Data Collection on Adverse Events of Anti-HIV
Drugs (DAD) study group reporting that ART was associated with a 26% increase in the
rate of myocardial infarction (MI) per year of exposure to ARV drugs. Islam et al. (2012)
also confirmed that the duration of exposure to ART might be one of the main factors
associated with risk of developing CVD.

These findings also suggest that melatonin supplementation does not alleviate this pro-
contractile effect of ART. This result contradicts results previously obtained in our labs,
where melatonin was shown to be a significant pro-relaxant (Smit-van Schalkwyk, 2016).
However, these inconsistent findings could be due to multiple factors. This study used a
higher dose of 10 mg/kg/day melatonin, where the previous study used a dose of 4
mg/kg/day. This study also used a treatment periods that were 2 weeks longer.
Furthermore, the previous study used melatonin to alleviate damage caused by nicotine
rather than ART, which may provide an indication that ART does not cause significant
enough damage in vivo for melatonin to act in a protective, vasodilatory manner. This
finding also correlates with the data obtained from the in vitro melatonin dose-response
studies, were we found that a shorter treatment period, with a lower melatonin
concentration was more protective.
Ach induced relaxations were similar for all treatment groups (Fig 5.2 B), which is surprising considering the few vascular reactivity studies that have been conducted, all showed significant findings in Ach-induced relaxation studies (Jiang et al., 2010; Jiang et al., 2006). Previous studies found that ART had an anti-relaxation effect, but as mentioned above, these studies did not use the same animal model or ART regime. The results found in this study may be may shed light on the fact that first line ART is less likely to induce ED than other previously studies ART regimes.

Results from pre-treatment with the NOS inhibitor, show that L-NAME elicited a strong pro-contractile effect (Fig 5.3 A) and reduced vasodilatory responses in all the groups (Fig 5.4 A). These experiments not only confirmed that the vascular isometric tension model was functional, but also showed the involvement of NOS and NOS-derived NO in anti-contractile and vasodilatory responses of the vasculature.

### 7.2.2 Endothelium-independent Aortic Ring Investigations

To determine whether differences in Ach-mediated relaxation were attributable to differential responses of the vascular smooth muscle to NO, relaxation responses to the NO donor SNP were evaluated.

For these endothelium-independent investigations, the Phe contraction studies expectedly followed the same trend as seen for the endothelium-dependent investigations, where cumulative contraction with Phe, showed that the both the ART and melatonin + ART treatment groups exhibited significant pro-contractile properties compared to the control group (Fig 5.5 A).

When SNP was used to induce relaxation, no significant differences where seen between any treatment groups (Fig 5.5 B). This finding indicates that the smooth muscle component of the NO-cGMP pathways remains unchanged during ART treatment (Sutliff et al., 2002). Further protein investigations should be conducted to examine whether this result was simply due to the fact that ART did not cause any oxidative damage, or whether SNP resulted in normal relaxation despite the presence of damaging increased levels of superoxide. This is a phenomenon that many studies have previously demonstrated as possible (Eberhardt et al., 2000; Gunnett, Heistad, & Faraci, 2002), and therefore should be further examined. Mechanisms for the dissociation between elevated superoxide and unaffected relaxation to SNP remain unclear (Sutliff et al., 2002).

Since endothelium-dependent and independent relaxation studies both showed no significant differences between groups, it is difficult to draw any conclusions as to whether
ART-induced damage is limited to the endothelium and endothelium-dependent factors like eNOS.

**7.2.3 Western Blot Analyses**

Western blot analyses were performed on snap-frozen aortic tissue of control, melatonin, ART and melatonin + ART treated animals from the endothelium-dependent studies, in order to determine whether the functional data of the vascular reactivity investigations could be explained by changes in intracellular signal transduction pathways.

No statistically significant changes were seen between any treatment groups in iNOS (Fig 5.7), nitrotyrosine (Fig 5.8) or p22 PHOX (Fig 5.9) expression, respectively. These results were unsurprising considering no differences were seen when Ach induced relaxation was examined. However, these results indicate that the pro-contractile influence elicited by ART is most likely not due to increased NADPH-dependent superoxide production or due to peroxynitrite-related damage.

**7.2.3.1 eNOS**

When the phosphorylated eNOS was expressed as a ratio of total, we saw that there were no significant differences between any treatment groups (Fig 5.6 C), despite that fact that eNOS phosphorylation was significantly lower in the melatonin + ART treatment group compared to the control (Fig 5.6 A). The P/T ratio obtained for the treatment groups was most likely due to the fact that the trends seen in eNOS phosphorylation and expression were relatively similar, notwithstanding that no significance was seen between groups for eNOS expression. This result was expected since there were no changes in vasodilatory capacity between groups during relaxation studies. This result does however shed more light on the mechanisms behind the ART-induced pro-contractile responses observed in this study. Previously it was proposed that this effect might be due to decrease NO production from decreased eNOS expression. These results however show that it is unlikely that decreased eNOS expression is the mechanism at work. This might point towards other contractor factors like endothelin-1 (ET-1), thromboxane A (TXA2) and angiotensin II (Cines et al., 1998; Mudau et al., 2012) being upregulated. However, further investigations would need to confirm this.
7.2.3.2 Cleaved Caspase-3

The ART and melatonin + ART combination treatment groups showed significantly decreased cleaved Caspase-3 expression compared to the control, while the ART treatment group also showed significantly decreased cleaved Caspase-3 expression when compared to the melatonin treatment group (Fig 5.10). These findings are very similar to those seen in the cell viability investigations of the in vitro study (Fig 4.17). Once again it seems that ART is the factor associated with decreased apoptosis. This finding further cements the anti-apoptotic role of ART, although in these in vivo studies, it would appear that this is not influenced by increased NO levels, since there was no increase in vasodilation seen in the ART treated groups. Other studies that looked at cleaved caspase-3 expression during ART have all been in the presences of HIV, and therefore it is difficult to differentiate between the effects of HIV and ART (Cook-Easterwood, Middaugh, Griffin, Khan, & Tyor, 2007; Pitrak et al., 2015).

7.2.4 Antioxidant Capacity Studies

The ORAC assay showed no significant differences in antioxidant capacity between treatment groups (Fig 5.11). As suggested for the in vitro studies in section 6.5, perhaps further antioxidant capacity assays should be performed to confirm this result.

7.3 Concluding Remarks

Novel ex vivo studies examining the acute effects of melatonin and ART treatment indicated that ART elicited a pro-contractile effect when administered to a healthy rat aorta. This resulted in the ART treatment group being pre-conditioned and consequently contracting less when compared to all other treatment group. It was also established that melatonin does not play a protective role in the presence of ART this situation, since the melatonin + ART treatment group relaxed significantly less than the control group. These findings are very interesting and need further elucidation through protein investigations.

In vivo studies demonstrated that ART has no effect on endothelium-dependent or independent relaxation, as shown by the lack of differences seen between groups in both Ach- and SNP-induced relaxation studies. Both these studies did, however, show that ART seems to have a pro-contractile effect. Further investigations are needed to establish by what mechanism ART elicits a pro-contractile response, yet does not affect vasorelaxation.
Protein investigations were relatively in line with these findings, with cleaved caspase-3 investigations shedding more light on the fact that NO levels are most likely not involved in the anti-apoptotic effects of ART.
8 – Overall Conclusions

This study aimed to investigate the effects of melatonin supplementation during first line FDC ART, through an in vitro, ex vivo and in vivo approach.

The in vitro dose-response studies showed that high concentration ART has the ability to induce modest increases in NO and RNS production. Combination studies highlighted that melatonin has the ability to ameliorate necrotic damage caused by ART in AECs. Signalling investigations ruled out endothelium-dependant NOS as a source of the increased NO observed in dose-response studies, as well as excluded NADPH oxidase-dependent super oxide as the likely source of the increase in RNS observed. Finally, ART was shown to have an anti-apoptotic effect, which may have been facilitated by the increase in NO observed in the dose-response studies. Taken together, these results show that first line FDC ART is very mildly damaging to the endothelium, and can even be protective under certain circumstances. Melatonin was shown to be effective in increasing cell viability, but showed no promise of being protective in terms of decreasing excessive NO and RNS production. However, since excessive levels of NO and RNS were not observed or produced in this study, no conclusions can be drawn as to whether melatonin has the ability to decrease excessive levels of NO and RNS in vitro form the current findings.

Ex vivo and in vivo studies shed light on the fact that there are still many unknowns with regards to the underlying mechanisms involved in the pro-contractile effects of ART. Although ART was shown to definitely elicit contractile responses, the lack of findings in the relaxations studies makes it difficult to draw conclusions on the effects of ART on aortic endothelial health. Since ART was not shown to decrease relaxation capacity in any studies, it could be said that first line FDC ART did not induce endothelial dysfunction, since ART treated aortas showed sufficient vasodilatory responses when compared to untreated aortas. Signalling investigations further established that ART has an anti-apoptotic effect. Whether this finding is specific to the specific regime of ART used in this study, or a general characteristic of ART, would have to be further investigated. In in vitro studies, melatonin was also shown to not have a protective effect in contraction and relaxation studies, but this could perhaps just be said for the specific concentration and duration of treatment of melatonin used in this study. Ex vivo and in vivo studies overall showed that first line FDC ART does not induce reduced relaxation capacity, usually
associated with endothelial dysfunction, while 10 mg/kg/day melatonin treatment over an 8 week period did not provide protective effects.

Taken together, these *in vitro*, *ex vivo* and *in vivo* investigations allowed for a very broad overview of the topic and allowed for the examination of vascular endothelial function on a cellular and functional scale. Although it is difficult to fully compare the different models, each aspect of this study allowed for the development and finding of important investigations that still need to be conducted, which could eventually lead to a better understanding behind the mechanisms by which ART potentially causes endothelial damage.

Overall, from these studies, it cannot be explicitly said that South African first line FDC therapy induces damaging endothelial changes, and that melatonin has the ability to alleviate those changes. However, particular findings did indicate that further investigations are need to be conducted, specifically with regards to the necrotic effects of ART and the acute effects of ART.
8.1 Advantages, Limitations and Future Directions

Currently, very few studies exist that investigate the effects of South African first line FDC ART. This study also investigated the effects of ART in isolation from HIV infection, allowing for a more accurate perspective of the effects of ART. This study showed that ART induced necrosis as measured by propidium iodide, can be decreased by conjunctive melatonin treatment. This study also demonstrated that acutely administered ART caused short-term spontaneous contraction, which resulted in an anti-contractile response in ART-treated aortas when exposed to cumulative concentrations of Phe. Both these findings are novel and require further study to clarify the mechanisms by which these actions occur.

Limitations of this study include the use of only three fluorescent probes for the plate reader investigations. In particular, a marker for apoptosis and well as a more specific marker for mitochondrial ROS should be included. Antibodies specific to mitochondrial ROS could also be included to better understand the mechanisms of ART-induced ROS production.

Small sample sizes are a prominent limitation in the *in vitro* western blot analyses. A new method should be developed in order to allow for the combination of multiple parent cell lines when making a single set of lysates, where cells could potentially be treated and then frozen until enough different parent cell lines have been grown.

Another potential limitation of this *in vitro* study might be its potential to translate to clinical relevance, since the long-term effects of ART are the main cause of ART-induced ED, whereas a treatment period of only 24 hours was examined.

Time and equipment-constraints meant that L-NAME studies could not be conducted for the endothelium-independent SNP relaxation investigations. This would be an interesting aspect to add to these studies. Signalling investigations should also be conducted in future studies examining the endothelium-independent chronic effects of ART treatment, as well as the *ex vivo* acute effects of ART treatment.

8.2 Specific roles in the study

- All *in vitro* cell culture experiments were conducted in the Medical Physiology Division on the Faculty of Medicine and Health Science at Stellenbosch University, by the student, with the exception of the ORAC assay which was conducted by Dr D Blackhurst at the University of Cape Town.
- All animal drug treatments were prepared and administered by the student, with the exception of the ART gavaging procedure, which was performed by Mr N Markgraaff and Mr S Smit.
- Animal housing, care, feeding and treatment occurred in the Animal Housing Unit of the Faculty of Medicine and Health Science at Stellenbosch University.
- All ex vivo and in vivo experiments were conducted in the Medical Physiology Division on the Faculty of Medicine and Health Science at Stellenbosch University, by student, with the exception of the ORAC assay, which was conducted by Dr D Blackhurst at the University of Cape Town.

8.3 Research outputs associated with the study

- Rawstorne, J., Strijdom, H., Webster, I., Westcott, C. The effects of melatonin supplementation on vascular tissue during first line ART: an in vitro and in vivo study. 45th conference of the Physiology Society of Southern Africa, hosted by university of Pretoria, 27- 31 August 2017. (Received an honorary mention for one of the top presentations presented for the Wyndham Competition)
Appendix A: *In vitro* melatonin and vehicle concentration calculations

**Melatonin Preparation (1 nm; 1 µM; 10 µM):**

- Weight off 23.2 mg melatonin and add it to 150 µl methanol (MeOH).
- Vortex solution well and make up to 10 ml by adding 9.850 ml distilled water (dH₂O) = 10 mM Melatonin Stock
- To make up 10 µM melatonin: Add 10 µl of the 10 mM Melatonin Stock to 9.990 ml of medium.
- To make up 1 µM melatonin: Add 2 µl of the 10 mM Melatonin Stock to 19.998 ml of medium.
- To make up 1 nM melatonin:
  - Add 1 ml of the 10 mM Melatonin Stock to 9 ml of dH₂O = STOCK A (1 mM melatonin).
  - Add 1 ml of STOCK A to 9 ml of dH₂O = STOCK B (100 µM melatonin).
  - Add 1 ml of STOCK B to 9 ml of dH₂O = STOCK C (10 µM melatonin).
  - Add 2 µl of STOCK C (10 µM melatonin) to 19.998 ml of medium.

**MeOH Vehicle Preparation (1 nm; 1 µM; 10 µM):**

- Add 150 µl MeOH to adding 9.850 ml distilled water (dH₂O) = 10 mM Vehicle Stock
- To make up 10 µM vehicle: Add 10 µl of the 10 mM Vehicle Stock to 9.990 ml of medium.
- To make up 1 µM vehicle: Add 2 µl of the 10 mM Vehicle Stock to 19.998 ml of medium.
- To make up 1 nM vehicle:
  - Add 1 ml of the 10 mM Vehicle Stock to 9 ml of dH₂O = STOCK A (1 mM vehicle).
  - Add 1 ml of STOCK A to 9 ml of dH₂O = STOCK B (100 µM vehicle).
  - Add 1 ml of STOCK B to 9 ml of dH₂O = STOCK C (10 µM vehicle).
  - Add 2 µl of STOCK C (10 µM vehicle) to 19.998 ml of medium.

*All melatonin preparations are done in the dark*
Appendix B: *In vitro* ART and vehicle concentration calculations

Efavirenz (5.0 µM; 8 µM; 12 µM):

MW: 315.67 g/L

Dissolve 10mg in 1.5 ml of Methanol.

\[ C = \frac{m}{MV} \]

\[ = \frac{0.01g}{(315.67 \text{ g/l} \times 0.0015l)} \]

\[ = 0.010559551 \text{ M} \]

\[ = 21119 \mu\text{M} \]

5.0 µM:

\[ C_1V_1 = C_2V_2 \]

\[ 21119 \mu\text{M} \times V_1 = 5.0 \mu\text{M} \times 1000\mu\text{l} \]

\[ V_1 = 0.24 \mu\text{L per 1 ml medium} \]

8 µM:

\[ C_1V_1 = C_2V_2 \]

\[ 21119 \mu\text{M} \times V_1 = 8 \mu\text{M} \times 1000\mu\text{l} \]

\[ V_1 = 0.38 \mu\text{L per 1 ml medium} \]

12 µM:

\[ C_1V_1 = C_2V_2 \]

\[ 21119 \mu\text{M} \times V_1 = 12 \mu\text{M} \times 1000\mu\text{l} \]

\[ V_1 = 0.57 \mu\text{L per 1 ml medium} \]
Emtricitabine (5 µM; 7.5 µM; 10 µM):

MW: 247.25 g/L

Dissolve 10mg in 4 ml of Milipore H₂O.

\[
C = \frac{m}{MV}
\]

\[
= \frac{0.01g}{(247.25 g/l \times 0.004l)}
\]

\[
= 0.0101112235 \text{ M}
\]

\[
= 10111.2 \text{ µM}
\]

5 µM:

\[
C_1V_1 = C_2V_2
\]

\[
10111.2 \text{ µM} \times V_1 = 5 \text{ µM} \times 1000\mu l
\]

\[
V_1 = 0.5 \text{ ul per 1 ml medium}
\]

7.5 µM:

\[
C_1V_1 = C_2V_2
\]

\[
10111.2 \text{ µM} \times V_1 = 7.5 \text{ µM} \times 1000\mu l
\]

\[
V_1 = 0.74 \text{ ul per 1 ml medium}
\]

10 µM:

\[
C_1V_1 = C_2V_2
\]

\[
10111.2 \text{ µM} \times V_1 = 10 \text{ µM} \times 1000\mu l
\]

\[
V_1 = 1.2 \text{ µl per 1 ml medium}
\]
Tenofvir (80 nM; 400 nM; 1 µM):

MW: 635.51 g/L

Dissolve 10mg in 40 ml of Milipore H₂O.

\[
C = \frac{m}{MV} = \frac{0.01g}{(635.51 \text{ g/l} \times 0.04l)} = 0.0003934 \text{ M} = 393.38 \text{ µM} \quad (\text{Stock A})
\]

80 nM:

Dilute stock of 393.38 µM:

Add 127.1 µl of Stock A to 872.9 µl of Millipore water (Stock B) (Now the concentration of Stock B is 50 µM)

\[
C_1V_1 = C_2V_2
\]

\[
50 \text{ µM} \times V_1 = 0.08 \text{ µM} \times 1000\mu l
\]

\[
V_1 = 1.6 \mu l \text{ per } 1 \text{ ml medium}
\]

400 nM (from stock A):

\[
C_1V_1 = C_2V_2
\]

\[
393.38 \text{ µM} \times V_1 = 0.4 \text{ µM} \times 1000\mu l
\]

\[
V_1 = 1.02 \mu l \text{ per } 1 \text{ ml medium}
\]

1 µM (from stock A):

\[
C_1V_1 = C_2V_2
\]

\[
393.38 \text{ µM} \times V_1 = 1 \text{ µM} \times 1000\mu l
\]

\[
V_1 = 2.54 \mu l \text{ per } 1 \text{ ml medium}
\]
Appendix C: Western Blot and ORAC 1 nM melatonin, high concentration ART and vehicle concentration calculations (combined vehicles)

Control Group (Combined Vehicles):

- 1.14 µl MeOH per 1 ml medium (Melatonin & EFV vehicle)
- 3.74 µl (1.2 µl + 2.54 µl) dH₂O per 1 ml medium (FTC and TDF vehicles)
- 1000 µl medium – 4.88 µl (1.14 µl + 3.74 µl) = 995.1 µl medium

Melatonin Group (+Combined Vehicles):

- Weigh off 0.002 g melatonin and add to 10 ml MeOH = Melatonin/MeOH STOCK A
- Add 10 µl of STOCK A to 9.990 ml MeOH = Melatonin/MeOH STOCK B
- 1.14 µl Melatonin/MeOH STOCK B per 1 ml medium (Melatonin & EFV vehicle)
- 3.74 µl (1.2 µl + 2.54 µl) dH₂O per 1 ml medium (FTC and TDF vehicles)
- 1000 µl medium – 4.88 µl (1.14 µl + 3.74 µl) = 995.1 µl medium

ART Group (+Combined Vehicles):

*The same pre-made ART stocks were used that were mentioned in Appendix B:
EFV = 21119 µM (0.57 µl per 1 ml medium)
FTC = 10111.2 µM (1.2 µl per 1 ml medium)
TDF = 393.38 µM (2.54 µl per 1 ml medium)

- 0.57 µl EFV/MeOH per 1 ml medium (EFV vehicle)
- 1.2 µl FTC/dH₂O per 1 ml medium (FTC vehicle)
- 2.54 µl TDF/dH₂O per 1 ml medium (TDF vehicle)
- 0.57 µl MeOH per 1 ml medium (melatonin vehicle)
- 1000 µl medium – 4.11 µl (0.57 µl + 3.54 µl) = 995.9 µl medium

Melatonin + ART Group (+Combined Vehicles):

- Weigh off 0.004 g melatonin and add to 10 ml MeOH = Melatonin/MeOH STOCK A
- Add 10 µl of STOCK A to 9.990 ml MeOH = Melatonin/MeOH STOCK B
- 0.57 µl Melatonin/MeOH STOCK B per 1 ml medium (Melatonin vehicle)
- 0.57 µl EFV/MeOH per 1 ml medium (EFV vehicle)
- 1.2 µl FTC/dH₂O per 1 ml medium (FTC vehicle)
- 2.54 µl TDF/dH₂O per 1 ml medium (TDF vehicle)
- 1000 µl medium – 4.88 µl (1.14 µl + 3.74 µl) = 995.1 µl medium
Appendix D: Western Blot Raw Data Normalisation

1. “Volume Intensity” values are retrieved from Image Lab™- 5 Software for both the loading-control and the specific antibody (eNOS in the example below).

2. For each sample within a treatment group (which represents a single lane on the Western blot gel/membrane), the volume intensity of the specific antibody is divided by the volume intensity of the corresponding lane or sample of the loading-control. Eg: eNOS “Control group/lane 1” is divided by Loading-control “Control group/lane 1”

3. An average is then determined for each treatment group.

4. The data is then normalised by dividing the antibody/loading-control value of each lane by the average “Control group” antibody/loading-control value (see value highlighted in red).

5. An average value is the determined per treatment group.

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Stellenbosch University  https://scholar.sun.ac.za
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*In the case of eNOS, a P/T ratio also needs to be determined. “P/T ratio” refers to the phospho-eNOS volume intensity divided by the total-eNOS volume intensity for each sample. See pg 146 for example data.*
### Table: p-eNOS

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Appendix E: Ex vivo 10 µM melatonin, high concentration ART and vehicle concentration calculations (combined vehicles)

*Calculations made for 20 ml organ bath filled with Krebs Henseleit buffer (KHB)

Control Group (Combined Vehicles):

- 22.8 µl MeOH in 20 ml organ bath (Melatonin & EFV vehicle)
- 70.6 µl dH₂O in 20 ml organ bath (FTC and TDF vehicles)

Melatonin Group (+Combined Vehicles):

10 µM melatonin:

*For high concentration EFV (12 µM), 11.4 µl MeOH will be used in 20 ml system since EFV has a set stock concentration that cannot be altered. Therefore 11.4 µl MeOH needs to be used for final 10µM melatonin when in combination with ART (and EFVs MeOH vehicle). For melatonin alone, 22.8 µl MeOH needs to be used to account for EFV vehicle.

\[ C1V1 = C2V2 \]

\[ (C1) \times (22.8 \, \mu l) = (10 \, \mu M) \times (20000 \, \mu l) \]

\[ C1 = 0.00877193 \, M \]

MW: 232.278 g/mol

\[ m = (0.00877193 \, M) \times (232.278 \times 0.003) \]

\[ = 0.006 \, g \text{ melatonin in 3 ml MeOH} \]

- 22.8 µl Melatonin/MeOH in 20 ml organ bath (Melatonin & EFV vehicle)
- 70.6 µl dH₂O in 20 ml organ bath (FTC and TDF vehicles)

ART Group (+Combined Vehicles):

*The same pre-made ART stocks were used that were mentioned in Appendix B & C:

EFV = 21119 µM (11.4 µl per 20 ml medium)
FTC = 10111.2 µM (19.8 µl per 20 ml medium)
TDF = 393.38 µM (50.8 µl per 20 ml medium)

- 11.4 µl EFV/MeOH in 20 ml organ bath
- 19.8 µl FTC/dH₂O in 20 ml organ bath
- 50.8 µl TDF/dH₂O in 20 ml organ bath
- 11.4 µl MeOH per 1 ml medium (melatonin vehicle)
Melatonin + ART Group (+Combined Vehicles):

\[ C1V1 = C2V2 \]

\[ (C1) \times (11.4 \, \mu l) = (10 \, \mu M) \times (20000\mu l) \]

\[ C1 = 0.01754386 \, \text{M} \]

MW: 232.278 g/mol

\[ m = (0.01754386 \, \text{M}) \times (232.278 \times 0.003) \]

\[ = 0.012 \, \text{g melatonin in 3 ml MeOH} \]

- 11.4 \, \mu l \text{ Melatonin/MeOH in 20 ml organ bath}
- 11.4 \, \mu l \text{ EFV/MeOH in 20 ml organ bath}
- 19.8 \, \mu l \text{ FTC/dH}_2\text{O in 20 ml organ bath}
- 50.8 \, \mu l \text{ TDF/dH}_2\text{O in 20 ml organ bath}
Appendix F: In vitro melatonin stock preparation

*Calculations for 1 cage*

The rats get 10 mg/kg/day Melatonin which is calculated as follows:

10mg/kg/day = 10mg/1000g/day = 0.01mg/g/day

1. Weigh the rats on Monday mornings, and then calculate the average weight per cage

Example:

\[(293.1g + 283.6g + 312.4g + 295.8g)/4 = 296.2g\]

2. Multiply the average weight by 0.01mg/g/day

Example:

\[296.2g \times 0.01 = 3mg\] melatonin per day [A]

3. The week before treatment started, rats were given 200ml of normal water on Monday around 16:30, and the next day at the same time, the left over water was measured to calculate how much water that cage drank in 24 hours. These 7 values are then averaged per cage, and divided by the amount of rats in the cage (usually 4) to get the average amount drank per rat in that cage.

Example (Cage 1):

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<th>Amount drank (ml)</th>
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<td>152</td>
</tr>
<tr>
<td>Sunday</td>
<td>200</td>
<td>22</td>
<td>178</td>
</tr>
</tbody>
</table>
\[
(172 + 100 + 120 + 132 + 122 + 152 + 178)/7 = 140\text{ml average water drank by cage 1 per day}
\]

\[
140\text{ml}/4 = 35\text{ml average water drank per rat in cage 1 per day} \ [B]
\]

4. Calculate grams melatonin needed to ensure each rat is getting 10mg/kg/day when drinking an average of 35ml day. For the stock, 500ml of melatonin stock is prepared on Monday, and give the rats in their bottle (200ml per day). The stock lasts two days. The extra 100ml in the bottle is in case of spillage, and gets thrown away if not used. On Wednesday a fresh 500ml stock is made and on Friday 1 x 250ml stock (For Friday) and 1 x 500ml stock (for Sat & Sun) is made.

**Example:**

Melatonin (mg) needed in 500ml "stock" = \([A]/([B]/500\text{ml}) = [0.03\text{mg}]/([35\text{ml}]/500\text{ml})
\]

\[
= 42.4\text{mg Melatonin}
\]

\[
= 0.042\text{g Melatonin}
\]

5. Melatonin is then weighed off and dissolved in 1ml methanol (Melatonin is very insoluble in water, but only 0.2% methanol of the total stock is used - therefore the potential effect of the methanol is considered negligible) and added to 500ml distilled water.
Appendix G: *In vitro* ART stock preparation

The daily dose (mg / kg) of each active ingredient for a human was calculated as follows:

\[
A_{ld} = \frac{A_{l}}{m_{t}}
\]

\(A_{ld} = \) Daily dose of active ingredient (mg / kg).
\(A_{l} = \) Active ingredient (mg).
\(m_{t} = \) Average body mass of a human (kg).

(8.6 mg / kg / day EFV; 2.9 mg / kg / day FTC; 4.3 mg / kg / day TDF)

Daily dosage per animal (mg / kg) was calculated according to the USA FDA guidelines using a conversion factor of X 6 to convert human to rat ratio:

\[
A_{l_{rat}} = 6(A_{ld})
\]

\(A_{l_{rat}} = \) Daily dose of active ingredient for a rat (mg / kg)
\(A_{ld} = \) Daily dose of active ingredient (mg / kg)

(EVF: 51.6 mg / kg / day; FTC: 17.4 mg / kg / day; TDF: 25.8 mg / kg / day).

Thus, for treated groups, the daily dose of active ingredients per animal needed was calculated weekly based on the average TBM of each group as follows:

\[
A_{ld_{r}} = A_{l_{rat}}(M_{avg})
\]

\(A_{ld_{r}} = \) Daily dose of active ingredient per rat (mg / kg).
\(A_{l_{rat}} = \) Active ingredient for a rat (mg).
\(M_{avg} = \) Average TBM of rats in a group (kg).

The mass of the 3 active ingredients combined in each tablet was 1100 mg (600 mg EFV + 200 mg FTC + 300 mg TDF). The actual % (m / m) of the active ingredients in each tablet was calculated according to the mass of the tablet as follows:

\[
A_{l\%} = \frac{A_{l}}{m_{T}}
\]

\(A_{l\%} = \) % Mass active ingredient in the tablet.
\(A_{l_{T}} = \) Total mass of active ingredient, 1100 g.
\(m_{T} = \) Total mass of the crushed tablet (g).
The amount (mg) of crushed powder of the tablet needed for a treated group with a certain average TBM (350 g per animal for example) was calculated as follow:

**EFV:AI_{rat} = 51.6 mg/kg/day**

\[
AI_{d/r} = AI_{rat}(M_{avg})
\]

\[
= (51.6 \text{ mg} / \text{ kg})(0.35 \text{ kg})
\]

\[
= 33.22 \text{ mg}
\]

*\(AI_{d/r}\) = Daily dose of active ingredient per rat (mg / kg).

*\(AI_{rat}\) = Active ingredient for a rat (mg).

*\(M_{avg}\) = Average TBM of rats in a group (kg).

If the % (m/m) of active ingredient is 68 % (for example), the amount (mg) of tablet powder weighed off was calculated as follow:

\[
AI_{w} = AI_{d/r}(AI_{%})
\]

\[
= 33.22 \text{ mg (68 %)}
\]

\[
= 22.58
\]

*\(AI_{w}\) = Mass of the tablet powder weighed off per rat (mg).

*\(AI_{d/r}\) = Daily dose of active ingredient per rat (mg / kg).

*\(AI_{%}\) = Mass % active ingredient in the tablet.

To prepare enough of the FDC for 4 rats for 8 days:

\[
M_{TP} = m_{p}(N)(d)
\]

*\(M_{TP}\) = Total mass of crushed powder per group per week (g).

*\(m_{p}\) = Mass of the crushed tablet needed per rat daily (mg).

*\(N\) = Number of rats in the group.

*\(d\) = Number of days
Odimune™ tablets and crushed in a pestle and mortar and weighed according to the above formula. ARVs were made weekly by dissolving the required weight in the appropriate amount of dH₂O is needed.
Appendix H: SNP concentration calculations

- A pestle and mortar was used to crush 1 small SNP crystal.
- 0.001g of SNP powder was weighed off from the crushed crystal and and add to 50ml 0.9% saline = SNP STOCK (60 µM). The tube was covered in tinfoil.

SNP Dilutions:

1.2nM:

\[ C1V1 = C2V2 \]
\[ (60 \, \mu M \times V1 = 0.0012 \, \mu M \times 25000 \, \mu l \]
\[ V1 = 0.5 \, \mu l \text{ in a 25 ml organ bath} \]

20nM

\[ C1V1 = C2V2 \]
\[ (60 \, \mu M \times V1 = 0.0188 \, \mu M \times 25000 \, \mu l \]
\[ V1 = 7.8 \, \mu l \text{ in a 25 ml organ bath} \]

70nM

\[ C1V1 = C2V2 \]
\[ (60 \, \mu M \times V1 = 0.0488 \, \mu M \times 25000 \, \mu l \]
\[ V1 = 20.3 \, \mu l \text{ in a 25 ml organ bath} \]

100nM

\[ C1V1 = C2V2 \]
\[ (60 \, \mu M \times V1 = 0.03 \, \mu M \times 25000 \, \mu l \]
\[ V1 = 12.5 \, \mu l \text{ in a 25 ml organ bath} \]

750nM

\[ C1V1 = C2V2 \]
\[ (60 \, \mu M \times V1 = 0.65 \, \mu M \times 25000 \, \mu l \]
\[ V1 = 270.8 \, \mu l \text{ in a 25 ml organ bath} \]
Appendix I: Male Wistar Rat Biometric Data

Increase in rat body mass (g) over the 8-week treatment period. Starting weights were recorded at week 0, and end weights were recorded before animal sacrifice.
Mean fluid intake (ml/rat/day) over the 8-week treatment period.
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