The Potential of *Spirulina platensis* to Ameliorate the Adverse Effects of Highly Active Antiretroviral Therapy (HAART)

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**Abstract:** The human immunodeficiency virus (HIV) is one of the most prevalent diseases globally. It is estimated that 37.7 million people are infected with HIV globally, and 8.2 million persons are infected with the virus in South Africa. The highly active antiretroviral therapy (HAART) involves combining various types of antiretroviral drugs that are dependent on the infected person’s viral load. HAART helps regulate the viral load and prevents its associated symptoms from progressing into acquired immune deficiency syndrome (AIDS). Despite its success in prolonging HIV-infected patients’ lifespans, the use of HAART promotes metabolic syndrome (MetS) through an inflammatory pathway, excess production of reactive oxygen species (ROS), and mitochondrial dysfunction. Interestingly, *Spirulina platensis* (SP), a blue-green microalgae commonly used as a traditional food by Mexican and African people, has been demonstrated to mitigate MetS by regulating oxidative and inflammatory pathways. SP is also a potent antioxidant that has been shown to exhibit immunological, anti-cancer, anti-inflammatory, anti-aging, anti-diabetic, antibacterial, and antiviral properties. This review is aimed at highlighting the biochemical mechanism of SP with a focus on studies linking SP to the inhibition of HIV, inflammation, and oxidative stress. Further, we propose SP as a potential supplement for HIV-infected persons on lifelong HAART.

**Keywords:** HAART/ARVs; *Spirulina platensis*; oxidative stress; HIV; antioxidant; inflammation; HAART toxicity; MetS

1. **Introduction**

The human immunodeficiency virus (HIV) has continued to be a global public concern due to its widespread infection rate and alarming mortality rate [1]. The Joint United Nations Programme on HIV/AIDS (UNAIDS), in its most recent report in November 2021, estimated that 37.7 million people globally are living with HIV. It was also reported that about 1.5 million new HIV infections and 680,000 AIDS-related deaths have occurred in the year 2020 [1–4]. In South Africa, approximately 8.2 million people were living with HIV in the year 2021 [4]. According to the South African mid-year population statistics 2021, there has been an unprecedented increase from 79,420 to 85,154 HIV-AIDS-related deaths in 2021 [4]. Recently, the easy availability of antiretrovirals (ARVs) has tremendously changed the pattern of death. ARVs have also helped prolong the lifespan of HIV-infected people in South Africa. Globally, about 27.5 million HIV-infected persons had access to ARVs in 2020, while approximately 5.6 million infected South Africans accessed ARVs in 2020 [1,4,5].

The highly active antiretroviral therapy (HAART) entails combining three or more antiretroviral drugs that are subject to the HIV-infected person’s viral load. HAART assists in regulating viral loads and preventing the progression to AIDS. Despite its success in prolonging HIV-infected patients’ lifespans, the use of HAART promotes metabolic syndrome (MetS) through an inflammatory pathway, excess production of reactive oxygen species (ROS), and mitochondrial dysfunction. Over three decades since its discovery, HAART has
significant improved the diagnosis and management of persons with HIV [6–12]. The persistence of MetS before and during HAART treatment further highlights the need for more studies on MetS inhibitory compounds, such as *Spirulina platensis* (SP) [13], *Moringa oleifera* [14,15], Curcumin [16], and Mangiferin [17]. However, there are other alternatives to combat MetS and these include exercise training [18], life style changes, and properly balanced healthy food choices [19,20].

SP is a blue-green microalgae commonly used as a traditional food by some Mexican and African people [21,22]. SP is mostly found in the alkaline water of volcanic lakes. In addition to its popular nutritional value, SP possesses various medicinal properties. It can induce both the humoral and cellular mechanisms of the immune system when consumed [22]. Interestingly, SP was linked with MetS-lowering properties, such as hypoglycemia [23], hypolipidemia [24], and hypotension [13]. Studies in rodents suggested that SP is particularly useful in preventing MetS [13]. SP contains oxidative stress inhibitors, phycocyanin and phycocyanobilin [25–27]. Previous studies have also demonstrated that SP inhibits oxidative stress [21,25,28,29] and promotes mitochondrial health [25,34]. Furthermore, SP can prevent the development of atherosclerosis [28] and diabetes [25]. The Food and Agriculture Organization (FAO) of the United Nations recommends *Spirulina* as a daily dietary supplement [35]. Microalgae including *Spirulina* are environmentally friendly and have a high rate of yield in large-scale production under controlled conditions [36]. *Spirulina*, in addition to its nutritional and medicinal security, has the potential to eliminate poverty. The considerable potential for sustainable financial development in a small-scale crop for nutritional enhancement was evident in China, where the production increase resulted in a dramatic increase in profit from USD 7.6 million to USD 16.6 million. *Spirulina* production is possible for small and marginal farmers as well as enthusiastic urban gardeners; this makes it easily accessible to the population [37]. Moreover, the health system can provide SP as medication to control adverse effects in people living with HIV on HAART with minimal costs.

Due to the increasing number of HIV-infected people and their high dependence on HAART, it is imperative to explore the anti-inflammatory and antioxidant mechanisms of SP against HIV and MetS. This review explores the anti-inflammatory and antioxidant mechanisms of SP in the inhibition of MetS and its potential as a supplement against HIV and ARV-induced MetS.

2. The Roles of HIV and HAART in MetS

HIV has often been associated with MetS [38], which results in cardiovascular diseases. Recent reports have linked some HIV-related features to MetS. These characteristic features include escalated cases of cardiovascular diseases, type 2 diabetes mellitus, dyslipidemia, immunodeficiency, high viral load, and atherosclerosis [7]. Recently, studies have suggested that HAART, in addition to the above-mentioned HIV-related features, actively induces MetS in persons with HIV [7,39–41]. Earlier studies by Palios et al. (2011) on arterial stiffness, displayed by pulse wave velocity (PWV), and markers of MetS, reported that persons with HIV exhibited an increased degree of PWV when compared with the healthy controls. Subsequently, persons on HAART have been shown to exhibit a similar PWV as persons with hypertension [39,40]. The prevalence of MetS in HIV-infected people receiving antiretroviral (ARV) treatment was higher when compared to the general population. This prevalence was attributed to age, physical inactivity, and a low cluster of differentiation 4 (CD4) count [42–44]. The patient response to HAART varies; some antiviral drugs can successfully suppress the plasma viral load without increasing the CD4 count; this allows the risks of opportunistic infections and abnormalities. Failure to increase the CD4 count during HAART may be due to several factors, including drug resistance, low CD4+ T-cell count at the initiation of HAART, the advanced stage of the disease, and a low adherence to HAART [45].

The nuclear factor-kappa-light chain-enhancer of the activated B cell (NF-κB) is a protein complex that is responsible for DNA and cytokine (IL-1β, IL-6, and TNF-α) tran-
scription, chemokine activation, and the survival of cells [7]. NF-κB is also a well-known mediator of inflammation that promotes HIV transcription [7,46].

Inflammation linked to HAART is evidenced by the persistently high levels of interleukin 6 (IL-6), C-reactive proteins, and D-dimers [47]. HIV creates chronic pro-inflammatory conditions that promote MetS [7]. Studies have shown that HIV is associated with inflammation, apoptosis, and mitochondrial dysfunction [7,48]; however, the mechanism that links HIV with MetS remains unclear [7]. Herein, we highlight the significance of SP as an anti-inflammatory supplement for HIV-infected people on lifelong HAART and its mechanism of inhibition on MetS.

3. Spirulina Species

*Spirulina* has three commonly investigated species due to their potential therapeutic nature and high nutritional content. These *Spirulina* species include *Spirulina platensis* (SP) (otherwise known as *Arthrospira platensis*), *Spirulina maxima* (*Arthrospira maxima*), and *Spirulina fusiformis* (*Arthrospira fusiformis*). These *Spirulina* species are also classified as oxygenic photosynthetic bacteria under Cyanobacteria and Prochlorales [49–54]. SP is found in alkaline water with abundant bicarbonate and saline [22,55]. *Spirulina* species are generally three-dimensional helix microstructures [56] protected by a cell wall composed of complex sugars and proteins [22]; however, helical transformation results after mature trichomes divide into hormogonia, binary fission, and undergo length elongation [57]. SP is considered an antioxidant and anti-inflammatory agent [58]. It is also considered as a nutraceutical food supplement due to its high content of proteins, vitamins, and minerals. Moreover, its composition includes chlorophyll, phycocyanin, and carotenoid. Chlorophyll has antioxidant and antimutagenic properties [59,60], carotenoids are vitally important antioxidants with cancer-inhibiting abilities [53], and phycocyanin is a Bili protein with antioxidant and radical scavenging properties [61]. Moreso, SP has also been credited for its cancer- and viral infection-suppressing abilities [62].

3.1. Role of *Spirulina* in the Inhibition of Oxidative Stress

Recently, research has unveiled the important roles of nicotinamide adenine dinucleotide phosphate (NADPH) oxidase in the production of reactive oxygen species (ROS). NADPH is contained in the nervous system and its assemblage and activation generate free radicals (FR) which subsequently destroy cells. *Spirulina* is a potent inhibitor of NADPH oxidase, which has been one of the proteins responsible for the production of ROS and subsequent oxidative stress [54,63,64]. SP has the potential to inhibit oxidative stress by blocking NADPH oxidase [21,25,28,29] and to enhance (Figure 1) mitochondrial health by promoting an antioxidant response [30–33]. Furthermore, SP prevents FR-induced apoptotic cell death through the inhibition of oxidative stress [65].

3.2. Multitargeted Therapeutic Roles of SP

SP has a therapeutic effect against vascular diseases, cancer, diabetes, neurodegenerative diseases, and inflammatory disorders [66]. *Spirulina* treatment enhances the NF-kappa B-directed luciferase expression [67]. It has antiallergic effects [54], including against allergic rhinitis in humans [68]. *Spirulina* is an immune booster [22]. It prevents cellular aging, infectious diseases, and promotes a strong immune system [57]. *Spirulina* has central neuroprotective effects in rodents [69]. It is also associated with the inhibitory effects against numerous viruses, such as HIV-1, herpes simplex virus 1 and 2 (HSV-1 and HSV-2), human cytomegalovirus (HCMV), influenza type A, measles, and other enveloped viruses [33,70–73]. Moreover, it has antimitagenic and anticancer effects [57]. *Spirulina* is an effective treatment against chronic arsenic poisoning with melanosis and keratosis [74]. SP shares similar chemical structures and physiological activities with bilirubin [25–28,75]. SP antioxidant properties are due to its composition and the presence of phycobilins, phycocyanin, and phycocyanobilin [25–27] (Figure 2). Phycobilins are similar in structure to bile pigments such as bilirubin, a well-known ROS scavenger [22,76]. Phycocyanin has
been proven to possess antioxidant and anti-inflammatory activities [34,54,63,64,77,78]. Phycocyanin is also structurally similar to biliverdin, a strong inhibitor of NADPH oxidase and inflammation-induced radicals [25,34]. A study conducted by Zheng et al. (2013) indicated that phycocyanin normalized urinary and renal oxidative stress markers and the expression of NADPH oxidase components. Furthermore, phycocyanobilin, bilirubin, and biliverdin inhibited NADPH-dependent superoxide production in renal mesangial cells [25]. The study also demonstrated that SP may be used in a therapeutic approach to prevent diabetic nephropathy through the inhibition of oxidative stress [25]. Thus far, phycocyanine, the most powerful natural antioxidant, is only present in cyanobacteria and thus in spirulina.

Figure 1. Diagrammatic representation of Spirulina reduction of oxidative stress via several pathways. Spirulina inhibits NADPH oxidase, reduces ROS, blocks FR-induced apoptosis, and promotes mitochondrial health (Created with BioRender.com, accessed on 17 November 2021).

Figure 2. The 2D chemical structures of phycocyanin, phycocyanobilin, bilirubin, and biliverdin (prepared by author using maestro 11.2).
SP exhibited neuroprotective activities through antioxidant and anti-inflammatory effects [79]. It has also shown significant antioxidant activity in vitro by scavenging nitric oxide and preventing DNA damage by scavenging hydroxyl radical (Figure 3) [80]. Antidiabetic and anti-inflammatory properties of SP [81] are based on its significant free-radical scavenging activities. SP contains compounds that fall under a broad spectrum of antioxidant agents, such as alkaloids, flavonoids [82], and phycocyanin [28,83–86]. Moreover, it provides trace minerals for the synthesis of antioxidant enzymes, demonstrated by the antidiabetic response in rats [87]. It has potential benefits in assisting with the reduction in chronic inflammatory conditions [88]. Spirulina incorporated into skin cream showed promising results as an anti-inflammatory and a wound-healing agent [89].

Figure 3. Antioxidant and anti-inflammatory effects of SP (created with BioRender.com, accessed on 17 November 2021).

Spirulina against HIV-1 demonstrated its ability as an antiviral compound. Studies have demonstrated the ability of SP in the inhibition of HIV-1 replication in human T-cell lines, peripheral blood mononuclear cells (PBMC), and Langerhans cells (LC). The inhibition of the viral production by spirulina extract (between 0.3 and 1.2 µg/mL) was found to be approximately 50% in PBMCs [57]. More studies are needed to fully understand the mechanism behind the inhibition of HIV by SP.

3.3. Mechanism of Action

SP contains several vital antioxidant and anti-inflammatory compounds as mentioned above, such as chlorophyll, phycocyanin, and carotenoids (β-carotene). The antioxidant and anti-inflammatory properties of phycocyanin have been determined in numerous studies [28,83–86,90–97]. Phycocyanin is responsible for reducing oxidative stress and NADPH oxidase [28]. It scavenges free radicals, such as alkoxy, hydroxyl, and peroxyl radicals, and decreases nitrite production and inducible nitric oxide synthase (iNOS) expression. Phycocyanin also inhibits liver microsomal lipid peroxidation [28,83–86,90–97].

Phycocyanin has been proven to inhibit the formation of the pro-inflammatory cytokine TNF-α and cyclooxygenase-2 (COX-2) expression. Additionally, it decreases prostaglandin E(2) production [28,83–86]. Phycocyanin prevents the degradation of cytosolic IκB-α, which suppresses the activation of nuclear factor-κB (NF-κB) [28]. Furthermore, the inhibitory activity of phycocyanin is associated with the suppression of TNF-α formation in the macrophages [86]. In addition, phycocyanin exerts regulatory effects on mitogen-activated protein kinase (MAPK) activation pathways, such as the p38, c-Jun N-terminal
kinase (JNK), and extracellular-signal-regulated kinase (ERK1/2) pathways [98–100]. The second compound of SP, carotenoids, β-carotene to be specific, is an antioxidant that has anti-carcinogenic, antioxidant, and anti-inflammatory activities [101–103]. As a membrane antioxidant, β-carotene protects against singlet oxygen-mediated lipid peroxidation [101]. Beta carotene inhibits the production of nitric oxide and prostaglandin E(2) and suppresses the expression of inducible nitric oxide synthase (iNOS), COX-2, TNF-α, and IL-1β. The suppression of inflammatory mediators by β-carotene results from its' ability to inhibit NF-κB activation by preventing nuclear translocation of the NF-κB p65 subunit [102,104,105]. Studies have shown that β-carotene suppressed the transcription of inflammatory cytokines such as IL-1β, IL-6, and IL-12 in vitro [103]; this takes place in the macrophages. The third compound of *spirulina*, chlorophyll, can perform antioxidant and antimutagenic activities [59,60]. *Spirulina*'s mechanism of action is a concert of compounds, but it is not clear whether they all act simultaneously during demanding events.

SP promotes the activation and expression of heme oxygenase 1 (HO-1) and endothelial nitric oxide synthase (eNOS) [104,105]. HO-1 is suggested to play an important part in the adaptive reprogramming which could result in Nrf-2 activation, but this pathway is unclear, and more studies are required. Moreso, SP causes the activation of the Nrf2/HO-1 pathway [106]. Nrf-2 activation by SP results in the production and increased expression of antioxidant enzymes, such as superoxide dismutase (SOD) and catalase (CAT) [100].

### 4. Common Highly Active Antiretroviral Therapy (HAART) Combinations

HAART has several classes: nucleoside/nucleotide reverse transcriptase inhibitors (NRTIs), non-nucleoside reverse transcriptase inhibitors (NNRTIs) [48,107–114], protease inhibitors (PIs) [114–117], integrase strand transfer inhibitors (INSTIs) [118–123], fusion inhibitors (FIs) [124], and chemokine receptor antagonists (CCR5 antagonists) [125–127]. HAART is a specifically selected combination of NRTI and NNRTI, PI, or INSTI drugs responsible for the inhibition of viral replication by multiple virus targets [113,128–136]. However, HAART can cause adverse drug reactions. Furthermore, 2′,3′-dideoxy-3′-thiacytidine (3TC), 2′,3′-dideoxy-5-fluoro-3′-thiacytidine (FTC), TDF (Tenofovir Disoproxil Fumarate), ZDV (Zidovudine), and d4T (Stavudine) are associated with mitochondrial toxicity and oxidative stress [137–140]. Additionally, NRTIs are linked to toxic hepatitis, and PIs are implicated in insulin resistance and hyperlipidemia [114]. Chronic side effects linked to HAART include ROS-induced insulin resistance [141,142], lipodystrophy, gastrointestinal disorders [143], and cardiovascular disease [11,144]. Primarily, HAART is used for the treatment and prevention of HIV-1. These primary functions of HAART are achieved by attacking different components of the virus lifecycle which ensures inhibition regardless of the virus being resistant to one of the drugs [113,128–136]. A combination of two NRTIs (mostly FTC and TDF) and one NNRTI (e.g., EFV; Efavirenz) is a more favorable choice due to the convenience to dose, effectiveness, and less toxic effects compared to other drug combinations [114,145,146]. A combination of three NRTIs is less effective than two NRTIs with an NNRTI [147]. The d4T/ddI combination is associated with high toxicity and hence it is not often recommended [148]. The most popular NRTIs are cytidine analogs (XTC), FTC, 3TC, and TDF, which form part of the first-line therapy [149–153]. FTC and 3TC are similar in chemical structures with different pharmacokinetic and pharmacodynamic properties, and they have required deoxynucleosides for HIV DNA synthesis. They undergo phosphorylation through intracellular kinases to become FTC 50-triphosphate (FTC-TP) and 3TC-TP; triphosphate metabolites with FTC-TP are more efficiently incorporated during HIV DNA synthesis than 3TC-TP [145,154–157]. Moreover, 3TC-TP has a shorter intracellular half-life compared to FTC-TP [155]. The TDF and FTC combination has a synergistic effect, increasing intracellular metabolites, and they are recommended for pre-exposure prophylaxis (PrEP) [155,158–161]. Contraindications (Table 1) on HAART treatment are antiretroviral medication-specific and can be overcome by a change in the HAART combination to suit the individuals own treatment [107,108,119–121,131,162–165] or the supplementation and treatment of the symptoms.
Table 1. HAART drugs mechanism and their adverse effects.

| HAART                      | Mechanism                                                                 | Example                                      | Adverse Effect                                                                                   |
|----------------------------|---------------------------------------------------------------------------|----------------------------------------------|--------------------------------------------------------------------------------------------------|
| Nucleoside/Nucleotide reverse transcriptase inhibitors (NRTIs) | NRTIs require intracellular phosphorylation via host enzymes before they can inhibit viral replication. These agents are nucleoside or nucleotide analogs with an absent hydroxyl at the 3’ end that are incorporated into the growing viral DNA strand. They competitively bind to reverse transcriptase and cause premature DNA chain termination as they inhibit 3’ to 5’ phosphodiester bond formation. | abacavir, didanosine, lamivudine, stavudine, tenofovir, emtricitabine, atazanavir, and zidovudine | Mitochondrial toxicity, bone marrow suppression, anemia, and lipodystrophy. NRTIs inhibit mitochondrial DNA polymerase. Tenofovir may cause kidney injury or decreased bone mineral density or osteoporosis. Abacavir is associated with a CD8-mediated hypersensitivity reaction in patients with the HLA-B*5701 mutation. Didanosine is associated with a high risk of pancreatitis and hepatomegaly. |
| Non-nucleoside reverse transcriptase inhibitors (NNRTIs) | NNRTIs bind to HIV reverse transcriptase at an allosteric, hydrophobic site. These agents cause a stereochemical change within reverse transcriptase, thus inhibiting nucleoside binding and inhibition of DNA polymerase. | delavirdine, efavirenz, nevirapine, rilpivirine | Temporary rashes but may progress to Stevens–Johnson’s syndrome. Hepatitis may progress to liver failure. Efavirenz may cause teratogenicity. Risk of neural tube defects. NNRTIs (mostly rilpivirine) may result in QT prolongation. Numerous interactions with hepatic cytochrome P450 enzymes. Efavirenz is linked to various psychiatric and CNS effects, including, but not limited to: vivid dreams, delusions, sleep disturbances, dizziness, headaches, increased suicidality, psychosis-like behavior, and mania. |
| Protease inhibitors (PIs) | PIs competitively inhibit the proteolytic cleavage of the gag/pol polyproteins in HIV-infected cells. These agents result in immature, noninfectious virions. PIs are administered with boosting agents such as ritonavir or cobicistat to patients that are failing their initial HAART combination. | atazanavir, darunavir, indinavir | Hepatotoxicity, insulin resistance, hyperglycemia, hyperlipidemia, lipodystrophy, and PR interval prolongation. Other PIs are inefficient and have a high resistance and increased risk of nephrolithiasis, hence indinavir and saquinavir are no longer used. |
| Integrase strand transfer inhibitors (INSTIs) | Integrase inhibitors bind viral integrase and prevent viral DNA from being incorporated into the host cell chromosome. | dolutegravir, elvitegravir, raltegravir | Some patients may experience dizziness, sleep disturbances, or depression. Raltegravir and dolutegravir can cause rhabdomyolysis and myopathy. Dolutegravir can block the secretion of creatinine and occasionally cause a decrease in the GFR. It can also have interactions with several medications, including those that inhibit/induce CYP3A4 enzymes, metformin, rifampin, and antiepileptics. |
| Fusion inhibitors (FIs) | Fusion inhibitors bind to the envelope glycoprotein gp41 and prevent viral fusion to the CD4 T-cells. | enfuvirtide | Enfuvirtide is generally well tolerated though some patients may experience injection site reactions. |
| Chemokine receptor antagonists (CCR5 antagonists) | CCR5 antagonists selectively and reversibly block entry into the CD4 T-cells by preventing interaction between CD4 cells and the gp120 subunit of the viral envelope glycoprotein. | maraviroc | Some patients may experience dizziness or skin rashes. There is a risk of hepatotoxicity with allergic features, high risk of hepatic dysfunction. Drug–drug interactions should be a consideration if patients are taking concurrent CYP3A4 inhibitors or inducers. |

Abbreviations: HLA—human leucocyte antigen; CNS—central nervous system; GFR—glomerular filtration rate.
5. Mechanism of HAART-Induced Oxidative Stress, Inflammation, and Mitochondrial Dysfunction

The exact mechanism of HAART-induced oxidative stress has not been completely explored; however, studies have demonstrated the link between HAART use and oxidative stress. HAART is linked with lipid metabolism dysfunction through the induction of peripheral lipodystrophy. Lipodystrophy results from the impaired cytoplasmic retinoic-acid binding protein type 1 (CRABP1)-mediated cis-9-retinoic acid stimulation of peroxisome proliferator-activated receptor type gamma (PPAR-γ), leading to impaired differentiation and increased apoptosis of peripheral adipocytes. HIV-1 protease-inhibitors further inhibit the cytochrome P450 3A-mediated synthesis of cis-9-retinoic acid, one of the key activators of PPAR-γ [166]. Insulin resistance occurs following impaired fat storage and lipid release [141,142,166], which impacts the oxidant profile. The depletion of ATP production and mitochondrial dysfunction [8,9], and the depletion of mitochondrial DNA [167–169], are some of the ways HAART induces oxidative stress. HIV increases oxidative stress, and HAART increases lipid oxidation, which amplifies the ROS imbalance leading to increased oxidative stress complications [10,170].

Chronic exposure to HAART induces increased oxidative stress in endothelial cells and mononuclear cell recruitment, which leads to cardiovascular diseases in HIV patients on ARV therapy [12]. Inducing oxidative stress is common for protease and reverse transcriptase inhibitors [171]. HAART drugs induce oxidative stress in various ways, which include inhibiting DNA pol-γ activity and leading to mitochondrial dysfunction, and also through the depletion of mitochondrial DNA [167–169]. Studies have shown that patients on HAART have abnormally high levels of free oxygen radicals in sera compared to untreated HIV patients and HIV-uninfected participants [10,11,168,170,172,173].

Adverse drug reactions vary; TDF and lopinavir cause acute and chronic renal dysfunction [174–178]. TDF inhibits mitochondrial DNA-polymerase gamma, hence leading to the impaired function of energy-dependent transporters [179,180]. NRTIs are associated with the inhibition of mitochondrial DNA polymerase, lactic acidosis, subcutaneous lipoatrophy, peripheral neuropathy, and pancreatitis. The level of mitochondrial toxicity depends on the drugs; it is low with 3TC, FTC, and TDF, followed by ZDV, and higher with d4T. NNRTIs are associated with life-threatening skin reactions and toxic hepatitis. PIs are associated with insulin resistance and hyperlipidemia [7,114].

6. Combined and Synergistic Therapeutic Actions of HAART and SP

Studies have shown that the possible therapeutic effects of antioxidants may provide strategies in suppressing oxidative stress-induced comorbidities that emerge with the use of HAART therapy in HIV-infected individuals [12]. The combination of HIV and HAART has been associated with increased oxidative stress and lipid peroxidation. Furthermore, HIV or HAART induces ROS by inducing NADPH oxidase [181,182]. Interestingly, SP is a potent antioxidant [26,27] with anti-inflammatory activities [34], which makes it a potential supplement in the mitigation of oxidative stress induced by HAART adverse drug reactions. Moreover, SP can inhibit NADPH oxidase which is considered one of the main sources of ROS and free radicals in HIV-infected persons on HAART [34,181,182], resulting in reduced oxidative stress [28]. Moreover, β-carotene from SP protects against singlet oxygen-mediated lipid peroxidation [101]. Among HAART complications, TDF and lopinavir cause acute and chronic renal dysfunction [174–178]. Herein, phycocyanin from SP can normalize urinary and renal oxidative stress markers and inhibit NADPH-dependent superoxide production in renal mesangial cells [25], ameliorating renal dysfunction. Lately, SP has been an effective therapeutic approach to preventing diabetic nephropathy through the inhibition of oxidative stress [25]. These properties indicate SP as a potential agent to mitigate renal dysfunction caused by HAART. As mentioned above, NRTIs can inhibit mitochondrial DNA polymerase [179,180]. Studies in vitro showed that SP can enhance cell nucleus enzyme function, repair DNA synthesis [57], and enhance mitochondrial health [30–33,80]. Mitochondrial toxicity presented as peripheral neuropathy and lactic
Nutrients 2022, 14, 3076

Acidosis can be corrected by SP through providing trace minerals for the synthesis of antioxidant enzymes [87] and reducing chronic inflammatory conditions [88].

NNRTIs are associated with life-threatening skin reactions and toxic hepatitis [114], these conditions may be ameliorated by SP. Phycocyanin from SP can inhibit liver microsomal lipid peroxidation [28,83–86,90–97], and hence reducing toxic hepatitis. Moreso, SP incorporated in skin creams showed promising results as an anti-inflammatory and a wound-healing agent [89]; this can be beneficial in the mitigation of NNRTI-induced skin reactions. PI therapy induces insulin resistance and hyperlipidemia [7]. Additionally, HAART may be associated with a higher risk of myocardial infarction [114,183,184]. SP has a therapeutic effect against vascular diseases, cancer, diabetes, and neurodegenerative diseases [66]. In addition, the Spirulina family has also shown central neuroprotective effects in rodents [69] and may exert its neuroprotective activities through antioxidant and anti-inflammatory effects [79]. Therefore, SP is a recommended antioxidant to use as a supplement; the list of benefits is evident. It also has antiallergic effects [34], prevents cellular aging and infectious diseases, and promotes a strong immune system [57]. Herein, promotion of a strong immune system by SP can help increase CD4 cell counts, lower HIV viral loads, and slow down the progression to AIDS. Moreover, SP prevents FR-induced apoptotic cell death [65]: this may help decrease apoptosis of peripheral adipocytes induced by HAART. Chemically, SP is a recommended source of proteins, vitamins, and minerals [57], important nutrients for individuals on the HAART program. Finally, SP can assist HAART in the inhibition of HIV-1 replication because it has been shown to inhibit viral production in PBMCs.

There has been a number of clinical studies to investigate whether SP improves the quality of life in HIV-infected individuals. Marcel (2011) reported that insulin sensitivity in HIV patients improved more when a spirulina nutritional supplement was used instead of soybean [185]. Another study demonstrated for the first time that spirulina improves antioxidant capacity in people living with HIV [186]. Spirulina supplementation combined with a qualitative balanced diet showed potential to inhibit lipid abnormalities [187], significantly increase CD4 cells, and reduce the viral load in HIV-infected antiretroviral-naive patients [187–189]. However, there is limited information on the investigation of SP confirming the mechanism of antioxidant and anti-inflammatory effects and the impact on the quality of life in the HIV-positive population taking HAART. Thus far, it has been shown that supplementation with Spirulina platensis could improve the immune status of HIV patients on ART and decrease inflammatory and pro-oxidant levels [190]. The development of more clinical studies to confirm the SP protective effect in this population will answer many questions. The recommended concentrations of SP for daily supplementation varies, as studies have successfully used 19 g [185], 5 g [186], and 10 g [190,191].

7. Conclusions

HIV continues to be a major global cause of mortality. Besides the therapeutic benefits of HAART in HIV treatment, HAART has been linked to numerous adverse drug reactions which include oxidative stress, inflammation, and the disruption of mitochondrial function. SP as an antioxidant, anti-inflammatory, anticancer, and nutritional supplement possesses various corrective properties against attacks from viruses and bacteria. The corrective health properties of SP are largely attributed to antioxidant pigments found in SP. These pigments include chlorophyll, carotenoids, and phycocyanin which facilitate antioxidant, anti-inflammatory, and anticancer properties. The corrective properties of SP indicated in this review highlight its potential in the mitigation of HAART adverse drug reactions and MetS. The SP supplement potential is also supported by its ability to assist HAART in the inhibition of HIV-1. This review highlighted the corrective properties of potent antioxidant SP potential as a supplement for individuals on lifelong HAART experiencing MetS. Furthermore, this review highlights the need for more studies on SP and HAART synergy.
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