

Review

Potential Synergistic Supplementation of NAD⁺ Promoting Compounds as a Strategy for Increasing Healthspan

Arastu Sharma^{1,2}, Sophie Chabloz², Elisabeth M. Roider^{3,4,5} and Collin Y. Ewald^{1,*}

¹ Laboratory of Extracellular Matrix Regeneration, Institute of Translational Medicine, Department of Health Sciences and Technology, ETH Zürich, Schwerzenbach CH-8603, Switzerland

² AVEA Life AG, , Bahnhofplatz, Zug-6300, Switzerland

³ Cutaneous Biology Research Center, Department of Dermatology, Massachusetts General Hospital, Harvard Medical School, Charlestown, MA 02129, USA

⁴ Department of Dermatology, University Hospital of Basel, Basel-4031, Switzerland

⁵ Maximon AG, Bahnhofplatz, Zug-6300, Switzerland

* Correspondence: collin-ewald@ethz.ch (CYE)

Abstract: Disrupted biological function, manifesting through the hallmarks of aging, poses as one of the largest threats to healthspan and risk of disease development, such as metabolic disorders, cardiovascular ailments, and neurodegeneration. In recent years, numerous geroprotectors, senolytics, and other nutraceuticals have emerged as potential disruptors of aging and may be viable interventions in the immediate state of human longevity science. In this review, we focus on the decrease of nicotinamide adenine dinucleotide (NAD⁺) with age and the supplementation of NAD⁺ precursors, such as nicotinamide mononucleotide (NMN) or nicotinamide riboside (NR), in combination with other geroprotective compounds to restore youthful NAD⁺ levels. Furthermore, these geroprotectors may enhance the efficacy of NMN supplementation while concurrently providing their own numerous health benefits. By analyzing the prevention of NAD⁺ degradation through inhibition of CD38 or supporting protective downstream agents of SIRT1, we provide a potential framework of the CD38/NAD⁺/SIRT1 axis through which geroprotectors may enhance the efficacy of NMN supplementation and reduce the risk of age-related diseases, thereby potentiating healthspan in humans.

Keywords: aging; longevity; supplements; geroprotectors; SIRT1; NAD⁺; NMN; resveratrol; nutraceuticals; age-related diseases; flavonoids; senolytics; healthspan

1. Introduction

The doubling of life expectancy in the last hundred years can be attributed to progressive leaps in health and clinical care in the human population. However, at the expense of longevity comes the increased risk of disease and other physical ailments. Being the main risk factor for ever-increasing incidence of neurodegeneration, cardiovascular disease, cancer, and other medical ailments, aging has come to the forefront as a new clinical target within itself, opting to serve as a preventative measure for developing disease [1]. Aging can be characterized as universal, intrinsic, progressive, and deleterious to the health of an individual, and a multimodal onslaught of the body's physiological processes may be the culprit, whether it be through environmental damage or a decline in endogenous protective mechanisms or pure breakdown of the physiological process due to simple wear and tear [2]. Numerous pathways also are implicated and may become at risk due to aging, such as neurodegenerative disorders [3][4]. The societal burden of these diseases and age-related pathologies is ever-compounding and must be addressed with novel interventions to improve the quality of life for the aging population. The conundrum of aging and the resulting risk of developing age-related diseases has the propensity to affect every individual, further compounded by other biological and environmental risk factors. Targeting the *Hallmarks of Aging* exists as a model which would allow for

interventions ameliorating these processes: genomic instability, telomere attrition, epigenetic alterations, loss of proteostasis, deregulated nutrient-sensing, mitochondrial dysfunction, cellular senescence, stem cell exhaustion, and altered intercellular communication. Each intervention is thought to focus on these hallmarks, or even have a multi-modal crossover effect to further ameliorate the biological processes of aging [5]. Appropriate, widespread, and easily executable interventions are a key factor in combating the newly defined epidemic of aging and age-related disorders. In the field of orthomolecular medicine, the use of supplementation as a therapeutic approach to ameliorate the process of aging is highly applicable when selecting and formulating geroprotective treatments. Longevity supplements have shown exciting potential for improving healthspan, and research into longevity supplements is one of the most rapidly advancing fields researched in aging science today, however, further human studies are warranted [6]. Human trials utilizing regenerative or senolytic compounds to target skin aging, in particular, have shown improvement in the biological response to environmental stressors (such as UV and oxidative damage), which are often implicated in the process of aging [7–9]. Age-related deficits in cognition are also arising as viable targets of longevity boosting and geroprotective supplementation. These deficits can be contributed to excessive neuroinflammation, oxidative damage, and metabolic disruption, and nutraceutical interventions, particularly ones targeting these numerous disruptive pathways, have shown promise in ameliorating cognitive age-related decline [10]. Administration of combinations of micronutrients and other antioxidant compounds has also shown promising results in promoting longevity and ameliorating accelerated aging in mice [11]. One compound in particular, nicotinamide adenine dinucleotide (NAD⁺), is relevant in many biological processes as a cofactor for crucial metabolic function. These sensitive pathways are interrelated with Sirtuins and PARP1 and may be disrupted by external factors or intrinsic breakdown due to aging, leading to an increased risk for many disorders and cognitive decline associated with aging and neurodegenerative disorders such as Alzheimer's and Parkinson's disease [12](Zhang et al., 2019). Multiple cohort studies spanning various age groups have shown significant decreases in NAD⁺ levels with age; this dysregulation draws attention to NAD⁺ as a potential target for age-related pathologies [13]. Model organisms have shown delayed muscle atrophy and improvement in neurodegenerative pathologies in response to NAD⁺ precursor administration [14], as well as restoration of metabolic function [15]. The clinically relevant evidence of targeting NAD⁺ to replenish endogenous stores and invigorate the NAD⁺ salvage pathway is compounding, raising the question of how NAD⁺ administration may be able to affect aging or age-related deficits in humans. Given the importance of maintaining NAD⁺ levels for health and the age-dependent decline, restoring NAD⁺ levels is an attractive and feasible approach to promote health during aging. Here, we review the NAD⁺ metabolism and the molecular interactions with other longevity-promoting pathways. Numerous studies have analysed the effects of oral NAD⁺ or precursor administration on certain disease states, muscle performance, metabolic processes, and other metrics, however, the longevity-promoting effect of NAD⁺ precursors administration can be further optimized and enhanced by supplementation of NAD⁺ enhancing and supporting compounds [16]. Longevity boosting potential of many drugs, such as rapamycin and metformin, are apparent, but they are not typically available in the consumer market: other ingredients show promise in the context of longevity, but require further extensive research on their synergistic effects on the NAD⁺/SIRT1/CD38 Axis [17]. Therefore, we will discuss other potent ingredients and nutraceuticals that may influence and modulate the NAD⁺ axis involving SIRT1 and CD38 in longevity biological mechanisms to offer further geroprotection by combating the aging process and additionally providing each of the nutraceutical's own health benefits.

2. NAD⁺ Homeostasis

Nicotinamide adenine dinucleotide (NAD⁺) is one of the primary coenzymes in metabolic reactions and is highly involved with numerous other processes, such as energy

expenditure, metabolic and stress adaptations, and circadian rhythm maintenance. NAD⁺ levels sharply decline with age, and this decline can be contributed to the activity of CD38, an enzyme responsible for the degradation of NAD⁺, disrupting the NAD⁺ pathways during the course of aging [18]. NAD⁺ homeostasis in the body is critical for optimal biological function. NAD⁺ consuming enzymes (converting NAD⁺ to nicotinamide (NAM)) occupy a specific role in biological aging pathology and are implicated as interventional targets for geroprotection, such as NAD-degrading enzyme CD38, the sirtuins (SIRT) deacetylases, poly [ADP-ribose] polymerase 1 (PARP1: involved in the DNA damage response (DDR)), and the neuronal degenerating and NAD⁺ draining factor SARM1 [19,20]. The NAD⁺ salvage pathway is crucial for recycling the metabolites of biochemical reactions to replenish NAD⁺ stores in the body, and NAD⁺ can also be synthesized *de novo* from nicotinic acid (NA), nicotinamide riboside (NR), and nicotinamide mononucleotide (NMN) [21].

NAD⁺, Sirtuins, and Longevity-promoting Pathway

Disruption of proper NAD⁺ and loss of protective sirtuin activity have emerged as the prime targets for NAD⁺-based interventions [22]. Administration of NAD⁺ precursors, namely NR and NMN, have been shown to ameliorate age-related NAD⁺ pathology, particularly in the context of age-related diseases [14,23,24]. Increased age has been associated with a decreased NAD⁺/NADH ratio in human plasma through the deterioration of NAD⁺ stores rather than an increase of NADH [13], and replenishment of NAD⁺ rescues mitochondrial regulatory function from NAD⁺ induced pseudohypoxic mitochondrial stress during aging [15].

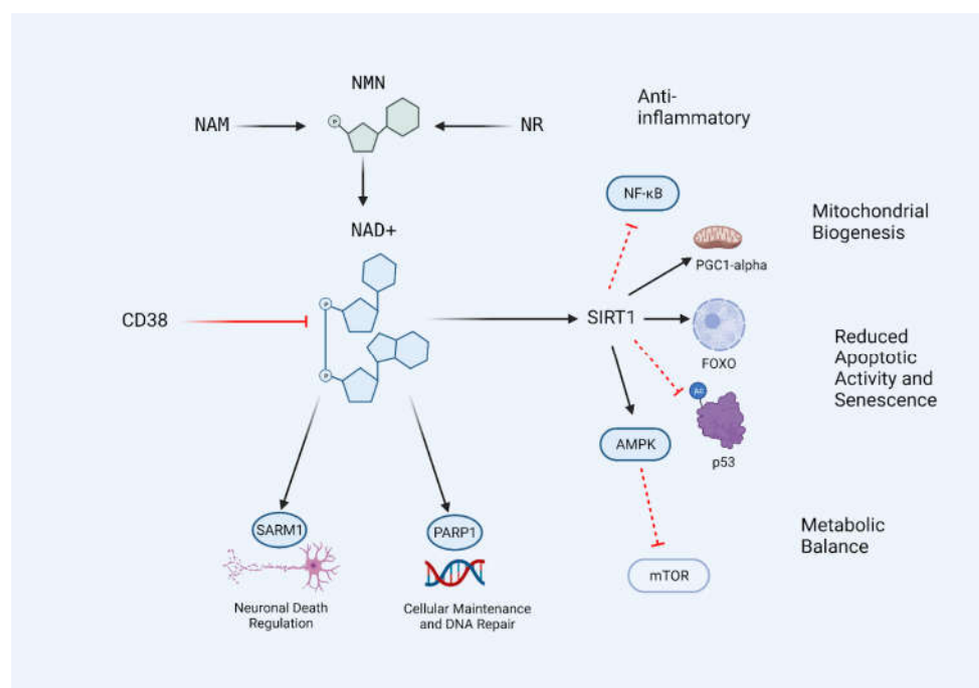


Figure 1. The CD38/NAD⁺/SIRT1 Axis. NAD⁺ levels in the body can be influenced by the supplementation of precursors nicotinamide (NAM), nicotinamide riboside (NR), and nicotinamide mononucleotide (NMN). NAD⁺ levels decrease with age, and are further metabolized by the activation of SIRT1, PARP1, SARM1, and CD38. Restoring NAD⁺ levels allows for an increase in SIRT1 activity due to increased substrate availability, resulting in the inhibition of age-promoting pathways and activation of adaptive and protective transcription factors and processes. The central lineage may be described as the CD38/NAD⁺/SIRT1 axis, and targeting this access with nutraceutical interventions may prevent the age-related decline of NAD⁺ levels in the body.

SIRT1 has been shown to be implicated in longevity, but experiments and analysis have yielded mixed results. For example, high-level athletes exhibit higher telomere

length and lowered insulin resistance, correlating with higher levels of SIRT1 expression [25]. SIRT1's beneficial activities may rely on the deacetylation and subsequent activation of Forkhead transcription factors FoxO and PGC1 α [26,27]. FoxOs is a transcription factor involved in stress resistance, cell cycle arrest, apoptosis, and tumour suppression, and activation of FoxOs has been linked with longevity in worms and flies [28,29]. PGC1 α exerts its influence on mitochondrial biogenesis, where deficits are apparent in metabolic disease states, and overexpression of PGC1 α has shown improved insulin sensitivity in muscle [30–32]. AMPK, involved in energy expenditure, is also exhibits strong similarity to and bi-directional innervation from SIRT1 and further inhibits mTOR, an inhibitory process that has also been linked to longevity, but further activates SIRT1 by increasing available NAD⁺ stores [33]. Furthermore, nuclear factor κ B (nf- κ B) signalling, a process involved in innate immunity, can be inhibited by SIRT1 activity to reduce prolonged inflammatory signalling [34]. Dependent on NAD⁺ stores in the body, SIRT1 activity arises as an interesting target in age-related pathway manipulation to promote longevity (Figure 1) [23,35–37]. The importance of proper NAD⁺ level maintenance for adequate SIRT1 activity during aging may be a key factor in regulating longevity.

NAD⁺ and Circadian Rhythm

NAD⁺ stores in the body are additionally extremely crucial for the programming of the circadian metabolic clock, where age-dependent NAD⁺ deficient mice experienced prolonged repression of CLOCK/BMAL1 transcription compared to younger mice with more abundant NAD⁺, resulting in disrupted and dampened mitochondrial and transcriptional oscillation [38]. NAD⁺ supplementation and restoration in circadian mutant mice have also been shown to re-establish proper respiratory oscillations and circadian metabolic regulation, particularly through SIRT3 regulatory activity [39]. A bountiful reserve of NAD⁺ and proper sirtuin activation are crucial for maintaining the integrity of the various endogenous clocks, and supplementation with NAD⁺ and precursors may potentially alleviate any age-related perturbations in these circadian processes [40]. NAD⁺ deficiencies are observed in these numerous age-related diseases, and NAD⁺-based interventions are currently underway in an attempt to ameliorate this common conundrum these diseases share [41–44].

CD38 and the Decline of NAD⁺

Intracellular levels of NAD⁺ are crucial for the proper function of several biological processes, including mitochondria metabolism [45]. In mitochondrial disease models, PARP inhibition or NR supplementation ameliorated disease deficits in metabolism and exercise capacity [46]. Strikingly, a cell surface glycoprotein, cluster of differentiation 38 (CD38), also acts as one of the main consumers of NAD⁺ stores in the body and is involved in immune activation and inflammatory signalling [47]. CD38, this NAD⁺ degrading enzyme, has also been shown to regulate SIRT1 activity along with NAD⁺ availability [48]. CD38 expression increases during aging, and CD38 knockout models exhibit heightened NAD⁺ levels and stark protection against obesity, metabolic disorders and cancer progression [49], potentiating CD38's role in the increased risk of developing age-related metabolic diseases [50]. However, CD38 deficiency may result in susceptibility to autoimmunity and decreased immune integrity, but further studies in humans are required to analyse the effects of insufficient CD38 [51]. Nevertheless, reduction of chronic inflammation is an aim of healthy aging; inflammatory profiles of senescent cells are associated with heightened CD38 activation and resulting NAD⁺ degradation through compounding processes of aging and inflammation [52]. Interestingly, this decline in NAD⁺ and increase in CD38 with age can potentially be attributed to another age-related phenomenon: the accumulation of senescent cells. Inflammation and aging intertwined to coin the term “inflammaging”, implying the build-up of inflammatory signalling due to cellular senescence, and notably the expression of the NAD⁺ degrading enzyme CD38 as a senescent cellular inflammatory profile. Senescent cells, through their senescence-associated secretory profiles

(SASP) of a milieu of IL-6, TNF- α , and CXCL 1 and 2, induce CD38 expression in M1 macrophages, causing NAD⁺ age-related decline in non-senescent phenotypic populations. These age-related senescent cells have been also shown to accumulate in liver and visceral white adipose tissue, and the resulting SASP induces pro-inflammatory M1 macrophage proliferation and enhanced CD38 expression [53–55]. The significant interplay between SASP, CD38, and NAD⁺ levels reveals an interventional target to further enhance available NAD⁺ in the body, particularly through strong inhibition of CD38 activity or expression well into the aging process to prevent any age-related biological deficits.

NMN as an NAD⁺ Boosting Therapeutic

When using an interventional method to target declining NAD⁺ levels, bioavailability and proper uptake of the proposed supplement must be analysed. NAD⁺ precursors, such as NR, NMN, and NAM, have been implicated in the proper uptake and subsequent biosynthesis of NAD⁺ [56,57]. For instance, NAD3®, an NR-based dietary supplement, was shown in another clinical trial in older adults to significantly improve NAD⁺ levels as well as improve blood lipid profiles, with a decrease in LDL:HDL cholesterol ratios [58]. A further meta-analysis revealed that NAD⁺ boosting mechanisms are successful in decreasing total cholesterol, triglyceride, and the LDL:HDL ratio [59]. Moreover, NMN in particular serves as a molecule of interest in the context of NAD⁺ support, mainly due to the fact that only one conversion step is required to reach NAD⁺ compared to other precursors. NMN supplementation has had numerous benefits, ranging from cardiovascular to neurodegenerative disease contexts, and proper uptake and resulting biosynthesis of NAD⁺ is observed in varying tissue types [60–62]. Successful increase in intracellular NAD⁺/NADH ratio was evident in a clinical trial in adults aged 40 to 65 over 60 days at a dosage of 300 mg, providing key safety and efficacy data to support the use of NMN Uthever™ as a valid interventional approach to increase NAD⁺ in a safe manner [63]. NMN supplementation has also been shown to target one of the hallmarks of aging, telomere attrition; short-term supplementation of NMN in pre-aging mice and humans, ages 45 to 60, showed a significant increase in telomere length in peripheral blood mononuclear cells, as well as altered faecal microbiota towards the benefit of immune and metabolic pathways [64], where specific administration in mice has been shown to increase the ratio of beneficial to harmful gut bacteria strains [65]. NMN supplementation has also been shown to enhance the miRNA vascular expression profile in aged mice [66]. Respiratory parameters and aerobic capacity can also be enhanced through 6 weeks of NMN supplementation in adult runners, likely attributed to the enhanced O₂ utilization occurring in skeletal muscle as a result of increased NAD⁺ availability [67]. Furthermore, 250 mg of daily NMN supplementation was shown to be well tolerated, efficacious when increasing NAD⁺/NADH ratio, and improved muscle function in aged but otherwise healthy men [68]. NMN supplementation has also been implicated in improving circadian parameters in older adults, where 12 weeks of 250mg daily NMN supplementation led to improvements in sleep quality, as measured through the PSQI, fatigue, and physical performance [69]. Having established the safety and efficacy of NMN supplementation to increase NAD⁺ levels in the body, numerous trials analyzing the effects of NMN on health, aging, and disease are constantly emerging.

A Combination Approach for Restoring Youthful NAD⁺ Levels During Aging

The momentum of clinical trials utilizing NAD⁺ boosting molecules is ever-increasing as interest in geroprotective trends grows. However, proper metrics evaluating the benefits and efficacy of intervention of NAD⁺ boosting supplements must be further standardized to offer significant and informative comparative effects of geroprotective interventions. NAD⁺ precursors have shown promising effects in human trials, but with the involvement of CD38 and SIRT1 pathways, many potential interactions exist to further amplify the NAD⁺ boosting effects of orthomolecular interventions. SIRT1 activation, NAD⁺ synergy, CD38 inhibition, and methylation support are all important factors when

considering maximizing the potential of NAD⁺ precursor administration. The synergistic effects and health benefits of compounds involved in these three processes will be further discussed to create a foundation to pave the way for potentiating supplement formulations for maximal NAD⁺ boosting effects. To maximize NAD⁺ boosting capacity, an elegant approach would be to supplement safe levels of NMN with other geroprotectors and nutraceuticals that will further enhance endogenous NAD⁺ levels, in this way ensuring the restoration of physiological levels of NAD⁺. Therefore, below we summarized different geroprotectors that enhance endogenous NAD⁺ levels but also have their own longevity benefits. For a selection of the following compounds, synergistic health effects of NMN and a geroprotector have been reported, suggesting the benefit of this combinatorial approach.

3. Potential Synergistic Nutraceutical Interventions

Stilbenes: Resveratrol and Pterostilbene

Some of the most extensively studied longevity-promoting nutraceuticals are the non-flavonoid phenolic stilbene compounds, resveratrol and pterostilbene [70]. These compounds are naturally occurring in grapes and berries and possess a plethora of highly anti-inflammatory and antioxidant characteristics, all the while combating many age-related disorders such as diabetes and cancer [71]. Clinical analyses have identified these stilbenes as safe and bioavailable substances with doses of resveratrol up to 5 grams and pterostilbene to 250 mg [72,73].

Resveratrol and pterostilbene have been shown to increase longevity [74] only in certain preclinical models, and pterostilbene has been shown to improve antioxidant function and increase longevity by activating longevity-associated genes, namely Sirt2, and has varying effects in male and female drosophila models with the same longevity inducing results but differing mechanisms [75]. Pterostilbene exhibits 80% increased bioavailability compared to resveratrol's 20% and possesses many of the same beneficial biologically modulating effects of resveratrol in the context of many disease states and aging by also upregulating the endogenous antioxidant enzymes such as superoxide dismutase (SOD) and glutathione reductase (GR) [76]. Pterostilbene's bioavailability becomes an important factor in modulating SIRT1 due to the high metabolism of resveratrol in the body and lower bioavailability; co-administration of resveratrol and pterostilbene may serve to increase the bioavailability and further compounding the benefits of each [77]. In response to particulate matter-induced oxidative damage, pterostilbene is able to mitigate the inflammatory response by decreasing mediators such as cyclooxygenase 2 (COX2), mitogen-activated protein kinase (MAPK), and the collagen degrading matrix metalloproteinase (MMP)-9 expression, preventing the keratinocyte aging processes [78–81]. Resveratrol provides further skin health benefits through its anti-angiogenic and wound-healing properties [82,83].

While resveratrol may also have numerous other targets, experimental models have shown a close association with the SIRT1 pathway when analyzing resveratrol dosage and supplementation as a study intervention [84,85]. Resveratrol has exhibited many health benefits but yielded mixed results when used for the treatment of disease specifically, and the role of resveratrol in the *direct* activation of SIRT1 has been heavily debated. Nevertheless, resveratrol seems to be at least indirectly involved in the activation of SIRT1 to induce the beneficial effects of caloric restriction, which has been shown time and time again to induce lifespan extension [86]. The activation of SIRT1 along with AMPK and PGC1 α in response to resveratrol is implicated, but despite this lack of clarity in interventional disease trials, resveratrol can still be considered as a SIRT1 and age-related metabolic modulator [87]. The benefits range from enhanced longevity to vascular improvements and motor function integrity into old age in mice.

Neuroprotection may also be enhanced through the administration of resveratrol, with a reduction of neuroinflammation, improved cognition, neuro-metabolic protection, and decreased risk of age-related neurodegeneration. resveratrol also possesses the ability

to improve oxidative stress in the vasculature by reducing superoxide generation and further increasing the antioxidant response of endothelial cells, dissipating age-related risk of cardiovascular disease, also initiated through the SIRT1-involved pathway. Anti-metastatic and anti-cancer properties are notoriously associated with resveratrol, as it has the ability to suppress tumour proliferation and unnecessary cellular division [88]. Interestingly, resveratrol has the ability to mimic caloric restriction (CR) in a similar fashion associated with the SIRT1 pathway to extend longevity, particularly through increased telomerase activity, FoxO3a, active regulator of Sirt1 (AROS), but conversely increased Hu antigen R (HuR) and decreased p53, which can be implicated in the prevention of cellular senescence but inhibition of tumour suppression abilities [89–91]. It has demonstrated the ability to improve biological parameters of kidney aging [92] and additionally improves arterial aging by reducing media thickness, inflammation, fibrosis, and oxidative stress, increased SIRT1, AMPK, and PGC1 α expression, SOD1 and 2, and decreased NADH oxidase 2 and 4 [93,94].

Cellular senescence can be inhibited by the administration of resveratrol at lower concentrations; however, at rather high concentrations, the proliferative potential of senescent cells is completely inhibited [95]. Resveratrol can even exert influence on reproductive aging, as seen in mice postovulatory oocytes. Rates of fertilization and blastocyst size increased in response to resveratrol administration, as well as upregulation of FoxO3a [96,97]. The cognitive effects of stilbene administration have gained more of a solid foundation in recent years, supporting its benefits for supplementation [98,99]. Cognitive impairment due to aging was attenuated by improved cognitive performance in older mice and decreased levels of inflammatory mediators such as IL-1 β and TNF- α in response to resveratrol [100], and aged mice that received resveratrol showed improvements in cerebrovascularity [101]. In response to a high-fat diet fed, resveratrol or alpha-ketoglutarate (aKG) improved brain and metabolic aging profiles in rats, notably through inflammatory response genes [102]. Witte et al., 2014, analysed resveratrol administration at 200 mg daily in older adults to affect memory performance, hippocampal connectivity, and glucose metabolism; memory retention increased in the interventional group along with increased functional connectivity in the frontal, parietal, and occipital regions of the hippocampus. Haemoglobin A1c (HB1Ac) and body fat decreased, implying lowered blood sugar, and the satiety hormone leptin increased, and both changes correlated with increased memory retention and increased functional connectivity. These changes were consistent with the benefits reported in previous studies, implicating the stilbene in enhanced cognitive function and improved metabolic profiles [103,104]. In the context of neurodegeneration and Alzheimer's disease patients with mild impairment, resveratrol treatment resulted in a decrease in neuroinflammatory profile and MMP9 expression, but further studies would also be required to validate the mechanism and efficacy in neurodegenerative disorders [105].

In the treatment of diabetes, resveratrol has been shown to improve oxidative and inflammatory microRNA profiles and reduce insulin resistance in diabetic humans [106]. Interestingly, resveratrol administration has been quite beneficial in ameliorating postmenopausal symptoms in older women; risk of osteoporosis and improvements of non-osteoporotic post-menopause-related bone density were reported in response to 75mg of daily resveratrol administration [107]. A 24-month crossover study in 125 postmenopausal women showed marked improvements in cognition, cerebrovascular function, and insulin sensitivity [108], and further sustained cognitive, cardio- and cerebrovascular benefits were reported over a 12-month study in response to the same daily dosage of 75 mg of resveratrol [109]. These results strongly implicate resveratrol in ameliorating age-related dysfunction in post-menopausal women as a preventative measure.

Resveratrol combined with senolytic flavonoid, quercetin, has been shown to also improve aging panels of the kidney in high glucose conditions by reducing glycation precursor, methylglyoxal, and increasing senescence marker protein 30 (SMP30) expression [110]. Rat heart tissue has also shown improved inflammatory, apoptotic, and oxidative panels in response to resveratrol administration, particularly through the reduction of the

inflammatory markers TNF- α and IL-gamma [111]. In particular, Ghanim et al. demonstrated that in response to a high-fat and high-carbohydrate meal, 100 mg of resveratrol and 75 mg grape skin polyphenols were able to upregulate oxidative stress-response transcription factor nuclear factor erythroid 2-related factor (Nrf2) binding activity to detoxification target genes NAD(P)H Quinone Dehydrogenase 1 (NQO-1) and glutathione-S-transferase pi 1 (GST-pi1), as well as ameliorating the effects of inflammatory mediators such as cluster of differentiation 14 (CD14), IL-1 β , and Toll-like receptor 4 (TLR4), potentiating the anti-diabetic, anti-inflammatory, and anti-atherosclerosis effects [85,112]. Induced cartilage stiffening in response to ribose administration can also further be ameliorated by resveratrol as a result of decreased glycation and advanced glycation end products (A.G.E.) because of resveratrol's and curcumin's competitive collagen II binding activity [113]. Resveratrol can further improve insulin sensitivity and skeletal aging through the SIRT1 pathway by reducing FoxO1-induced transactivation of pyruvate dehydrogenase kinase 4 (PDK4), implying an increase in muscular metabolism integrity [114,115].

As stated, resveratrol and its highly efficient and bioavailable sibling pterostilbene are key agents in combating the processes of aging and reducing the risk of developing age-related neurodegenerative, cardiovascular, cancer, and metabolic disorders. However, their role can be heightened when used in combination with NMN supplementation to produce optimal results as a cleverly tailored nutraceutical orthomolecular approach to delaying or even reversing the signs of aging [116] particularly when monitoring appropriate age-related biological markers indicating risk of these age-related diseases in response to resveratrol and pterostilbene administration [117,118].

While NAD⁺ is our compound of focus implicated in aging, resveratrol and pterostilbene have been shown to offer further benefits than the aforementioned by supporting NAD⁺ levels in the body. Coadministration of NMN and resveratrol resulted in increased NAD⁺ levels in heart and skeletal muscle compared to NMN alone, and NMN and ginsenoside coadministration resulted in higher NAD⁺ levels compared to NMN alone in brain, heart, kidney, and lung tissues, further supporting the dual supplementation approach to maximize NAD⁺ levels in the body [119]. Resveratrol also retains the ability to activate NMNAT1, the NAD⁺ synthetic enzyme, to increase NAD⁺ levels up to five times, providing a large substrate store for SIRT1 activation to promote longevity-enhancing pathways [120]. NR and pterostilbene co-administration additionally exhibit a 2-dose, dose-dependent increase in NAD⁺ levels in acute kidney injury patients [121], and repeated doses of NR and pterostilbene also exhibit an increase in NAD⁺ levels of 40% and 90% (double dose) in humans with a consistent increase over the 8-week trial duration [122]. resveratrol and quercetin are also able to promote cell protection and NAD⁺ levels along with NAD⁺ precursors by potentiating the cell rescue effects of said precursors in response to DNA damage [123].

The extensive benefits of resveratrol and pterostilbene warrant their consideration for inclusion in novel formulations of healthy aging interventions or supplements. Particularly noteworthy is the interplay between sirtuins, the NAD⁺ salvage/biosynthesis pathway, and genetic regulation of metabolic processes, which in turn may reduce the risk of age-related neurodegeneration, cardiovascular ailments, cancer, or metabolic disease. Resveratrol and pterostilbene are key agents in combating age-related and NAD⁺ deficiency-induced perturbations in biological and physiological machinery, and further enhancing the geroprotective effects of SIRT1 and NAD⁺ in the body by supporting the transient supplementation of NMN may serve as a crucial factor in maximizing the potential for this longevity based orthomolecular approach.

CoQ10

Coenzyme Q10 (CoQ10), or ubiquinol (oxidized form of ubiquinone) participates in the electron transport chain in the mitochondria, and low levels of CoQ10 are associated with diseases, such as neurodegenerative disorders, diabetes, cancer, fibrosis, and cardiovascular diseases [124]. Supplementation with CoQ10 in the context of disease states

aims to re-establish antioxidant activity to ameliorate the homeostatic disruptions [125]. CoQ10 has cardiovascular protective qualities and can result in improvements in hyperglycaemia, hypertension, reduced oxidative stress, and reduced risk of cardiac events [126]. Endogenous biosynthesis of CoQ10 sharply decreases with aging, and enhanced longevity can be seen with higher mitochondrial levels of CoQ10. Skeletal muscle integrity in elderly individuals is associated with higher plasma CoQ10 content, and inflammatory factors such as TNF- α , IL-6, and C-reactive protein (CRP) levels can be reduced as a result of supplementation [127]. It is also required for lipid integrity and protection of LDL oxidation against atherosclerosis [128]. Replenishing the age-related decline in levels of CoQ10 is imperative to reduce the risk of age-related diseases and lower the burden of age-inducing oxidative stress through nutraceutical supplementation [129]. CoQ10 supplementation, along with dietary interventions in elderly men and women, resulted in improved metabolic profiles, reducing metabolic and cardiovascular risk [130].

CoQ10 and NAD⁺ supplementation also portray a degree of synergy when analyzing maximum heart rate after exercise in chronic fatigue syndrome (CFS). Maximum heart rate was decreased and reports of fatigue improved [131]. CFS shares some characteristics with aging and age-deregulated pathways, such as inflammation and oxidative stress, and CoQ10 and NADH supplementation were further shown to improve fatigue and levels of NAD⁺/NADH, CoQ10, ATP, citrate synthase, and lowered levels of lipoperoxides [132].

The antioxidant, anti-inflammatory, and age-related effects of CoQ10 make it a highly appealing supplement to include in the comprehensive orthomolecular approach to battling the biological process of aging, particularly in the context of supporting NAD⁺ levels. Further studies are required to show, for instance, the beneficial synergistic effects of NAD⁺ precursor supplementations in combination with CoQ10.

Betaine

Trimethylglycine, known as betaine, was originally derived from the beetroot plant. It has osmoprotectant and anti-inflammatory properties. Betaine, along with methionine and choline, is one of the primary methyl group donors involved in DNA methylation, and DNA methylation rates have been directly correlated with available access to methyl donors [133]. Betaine also suppresses numerous inflammatory expression profiles, such as TNF- α , COX2, and NF- κ B activity [134]. The effects of betaine are implicated in postponing aging-related pathologies by promoting lipid and glucose metabolism, inhibiting inflammatory transcription processes, and reducing cellular ER stress [135].

Interestingly, betaine and choline levels are influenced by the degradation of the NAD⁺ precursors, namely NAM, which consumes more betaine than choline, potentially compromising the availability of a healthy pool of methyl donors [136]. The high burden of NAM degradation on betaine levels and epigenetic changes warrants adequate supplementation of methyl donors during concomitant NAD⁺ precursor administration, particularly NAM [137], but the effects of NMN or NAD⁺ conversion on methylation levels have not been explored. Concurrent supplementation of NMN, NAD⁺, or other precursors alongside betaine may prevent the proposed decline in betaine levels, maintaining proper methylation health and function.

Flavonoids: Quercetin, Fisetin, Luteolin/Luteolinidin, and Apigenin

Numerous flavonoids have also been shown to have a host of health benefits along with strong senolytic activity, such as fisetin, quercetin, luteolin/luteolinidin, and apigenin.

Fisetin and quercetin are potent anti-cancer agents and have been shown to have anti-tumour activity through Ca²⁺ induced tumour apoptosis along with improving cancer-related inflammatory profiles [138].

Fisetin has been shown to be a potent senolytic in older and progeroid models of mice, as well as in murine and human adipose tissue, and administration of fisetin improved lifespan and tissue homeostasis in mice [139]. Senolytic effect is a crucial

characteristic of an interventional dietary supplement, and fisetin has been shown to perform as one of the strongest among numerous other flavonoids and compounds. The safety and efficacy of fisetin are currently being investigated on reducing inflammation and improving walking speed in frail elderly individuals in randomized Phase 2 clinical trials (NCT03675724, NCT03430037). Interestingly, fisetin also portrays a degree of inter-relatedness to NAD⁺/NADH age-related pathology through its activation of SIRT1 by inducing mitochondrial fragmentation in a similar fashion to treatment with NAM and an increased NAD⁺/NADH ratio [140]. These findings point to the possibility that fisetin may have geroprotective effects in the context of the NAD⁺/SIRT1/CD38 pathway, but further experiments must be conducted to see measurable and explicit effects on the biochemical longevity axis.

Quercetin, another flavonoid with striking structural similarity to fisetin, is also considered a senolytic with many benefits in the realm of cardiovascular disease, neurodegeneration, inflammation, oxidative stress, cancer, and diabetes, and has been shown to be a geroprotective agent for *in vitro* models of premature aging [141,142]. Quercetin also exhibits attractive properties as a supplemental ingredient in the NAD⁺ boosting approach. Along with the previously mentioned apigenin, quercetin has also been shown in the same experiment to be a potent CD38 inhibitor to maintain the integrity of NAD⁺ stores and act as a protective agent against metabolic disorders [143]. Quercetin can potentially initiate SIRT1's intermediary action to result in anti-inflammatory and antioxidant effects in diet-induced atherosclerosis rat models [144]. NAD⁺/NADH balance is also further affected by quercetin's ability to oxidize NADH and decrease the NADH/NAD⁺ ratio, where larger concentrations of the flavonoid stimulated the Krebs's cycle, likely due to an increase in available NAD⁺ [145]. Nevertheless, quercetin earns a role in the modulation of the NAD⁺/SIRT1/CD38 axis due to its effects on altering the NAD⁺/NADH ratio, SIRT1 activation, and CD38 inhibition.

Luteolin, another flavonoid with similar health benefits as other geroprotectors, also has strong anti-inflammatory capabilities in relation to skin aging, skin disease, skin cancer, and cognition [146,147]. Luteolin is implicated in CD38 mechanisms along with apigenin and quercetin (and other compounds, such as luteolinidin and kuromanin) and has been shown to be a potent inhibitor of CD38, ultimately leading to an increase in available NAD⁺ [148]. Luteolinidin also exhibits CD38 inhibitory activity as well as myocardial protective attributes by stabilizing NADPH and mitigating the ischemic injury-induced upregulation of CD38 [149]. Flavonoids may further post longevity-promoting effects through the clearance of cellular senescence during concurrent administration of NAD⁺ supporting compounds, and further compounds may be tested on their activity on NAD⁺/NADH ratios and sirtuin activation [150].

The flavonoid apigenin is derived from plants, such as parsley or chamomile, and possesses highly anti-inflammatory, antioxidant, and anti-carcinogenic properties. It can reduce COX2, IL6, and TNF- α in response to the bacterial toxin lipopolysaccharide (LPS) [151]. Upregulation of antioxidant enzymes such as SOD, glutathione peroxidase (GPX), and glutathione reductase (GR) is also evident in response to lower dose apigenin administration [152]. Apigenin burnishes its anti-carcinogenic activity by downregulating key factors in cancer pathways and sensitizes tumour cells to chemotherapeutic interventions [153]. In the context of oesophageal cancer, IL-6 transcription is inhibited in response to apigenin as an intervention [154], and apigenin also exhibits anti-leukemic activity *in vivo* and in human cells by inducing apoptosis [155]. Apigenin can further attenuate metabolic complications leading to risk and characteristics of metabolic disorders, such as metabolic syndrome, atherosclerosis, and diabetes [156]. Apigenin reduces the expression of COX2 and can also produce anti-inflammatory effects in the context of cardiovascular disease, neurodegeneration, diabetes, and cancer [157]. Anti-obesity can also be an indication of apigenin supplementation, with marked reductions in visceral adipose tissue in high-fat-fed mice, and can preserve skeletal muscle in obese mouse models of obesity-induced skeletal muscle atrophy [158,159]. Apigenin also possesses the ability to ameliorate deleterious age-related changes in vascular endothelial function and structure due to

oxidative stress [160]. The anti-inflammatory and anti-cancer properties of apigenin make it appealing to utilize in the context of healthy aging and geroprotection [161]. However, the most notable property of apigenin in the context of NAD⁺ supplementation is the involvement with the SIRT1 NAD⁺ and CD38 axis. Apigenin administration increases the activation ratio of SIRT1, NAD⁺, NAD⁺/NADH and strongly inhibits CD38, thereby enhancing endogenous NAD⁺ levels. The resulting increase in SIRT1 reduced cellular senescence as a result of oxidative stress [162]. Apigenin is a potent inhibitor of CD38, and knockout models have shown resistance to metabolic disorders and exhibit increased NAD⁺ levels. Inhibition of CD38 in obesity models led to enhanced metabolic profile increased NAD⁺ levels, and decreased protein acetylation, possibly due to enhanced SIRT1 activity due to higher NAD⁺ availability [143]. CD38 inhibition through apigenin administration additionally restores mitochondrial function in response to oxidative stress and increases the NAD⁺/NADH ratio and SIRT3 activation [163]. Apigenin serves as a crucial component in the comprehensive approach to restoring age-related depletion of NAD⁺ levels through its strong inhibition of CD38, the degrading enzyme of NAD⁺. Through its CD38 inhibitory activity, apigenin becomes a valuable interventional agent when maximizing the effects of increasing NAD⁺ availability in the body through NMN supplementation.

Carotenoids: Astaxanthin and Lycopene

Astaxanthin is a potent antioxidant carotenoid compound that offers an array of health benefits by mitigating reactive oxygen species (ROS) and supporting mitochondrial integrity [164]. Unsurprisingly, astaxanthin is highly efficacious in utilizing SIRT1 to produce longevity-promoting effects in many experimental models. Astaxanthin was shown *in vivo* to soothe oxidative stress in response to brain injury by upregulating the expression of Nrf2 and SIRT1 while concurrently reducing the expression of pro-apoptotic factors, potentially reducing the risk of neuronal death events post-injury [165]. The effects of a high-fat-induced diet on cardiac and fibrotic damage can also be ameliorated by astaxanthin through upregulation of SIRT1, inhibition of inflammatory cellular mobility, and reduced incidence of collagen deposition leading to fibrosis post-injury [166,167]. Astaxanthin can additionally protect renal tissue post-injury through its upregulation of SIRT1 [168]. The most supportive finding of astaxanthin's ability to boost NAD⁺ availability was found by Zhang et al., where NMN, astaxanthin, and blood orange extract were utilized to successfully increase NAD⁺ in aging zebrafish, exhibiting superiority in raising NAD⁺ levels compared to supplementation of just NR, NR and astaxanthin, and even NR and pterostilbene [169]. This specific study can serve as a basis for future studies of supplements to hone in on effective and precise dosages and synergistic combinations to maximize the effect of NAD⁺ boosting approaches in humans.

Lycopene, another red-pigmented carotenoid, also possesses strong antioxidant and anti-inflammatory characteristics. Oxidative stress and the intrinsic aging process can lead to plasma lycopene depletion and impaired absorption, and supplementation has been shown to improve physical performance, osteoporosis, and skin aging [170]. Interestingly, lycopene activates SIRT1 to impart muscle angiogenesis and reverse insulin resistance in models of age-related vascular decline [171]. *In vivo* and *in vitro* models of D-galactose-induced aging were ameliorated in a superiority trial with astounding results in response to NMN and lycopene combination therapy, with increases in SOD; GSH-px, GPX, CAT, T-AOC, and evidence of senolytic abilities. Nrf2 was upregulated, and *in vivo* models showed enhanced cognition compared to NMN alone [172].

Curcumin

Curcumin has earned a role as a potent senolytic, along with the previously mentioned flavonoids. It has been implicated in improving senescence in age-related pathologies and has even exhibited modulatory effects on longevity-related pathways, such as the mTOR and FoxO [173]. In the context of neurodegeneration, curcumin has been shown

to upregulate SIRT1, and in the cardiovascular context, additionally AMPK [174]. Anti-cancer, migration, and anti-angiogenesis activity can be observed in response to curcumin administration in head and neck squamous cell carcinoma experimental cell models [175], and 6-week curcumin supplementation has been shown to improve antioxidant and aerobic capacity in human runners along with an increase in SIRT3 [176]. The connection between curcumin and sirtuins is evident, but the utilization of curcumin along with NAD⁺ boosting molecules must be first demonstrated in future superiority trials involving combination therapies.

aKG

Alpha-ketoglutarate (aKG) is a crucial metabolic intermediate that is involved in the Krebs's cycle and is implicated in the process of aging [177]. Along with other geroprotectors that inhibit the TOR pathway, AKG is able to additionally inhibit ATP synthase to extend lifespan in *C. elegans* in a fashion similar to caloric restriction [178]. Offering metabolic and antioxidant benefits [179], aKG produces lifespan-extending benefits which are additionally apparent in mice, and recent pilot clinical trials in humans have shown that Rejuvant, a novel formulation of aKG, was successful in reducing the biological age of study participants [180,181], however, data involving aKG and NAD⁺ interplay is lacking and must be explored further.

EGCG

The polyphenol epigallocatechin gallate (EGCG), found in green tea, has been touted to retain neuroprotective, antioxidant, and anti-inflammatory effects and is currently being analysed for its role in ameliorating numerous different diseases [182]. It has been shown to increase rat lifespan in response to oxidative stress [183]. However, the role of EGCG in SIRT1 modulation is unclear, as certain studies see upregulation of SIRT1 in response to administration [184], but others observe downregulation, particularly in cancerous cells [185,186]. EGCG may behave differently on SIRT1 depending on the context and which longevity factors must be upregulated to mitigate oxidative stress, and the effects of EGCG on NAD⁺/NADH ratio must be further observed.

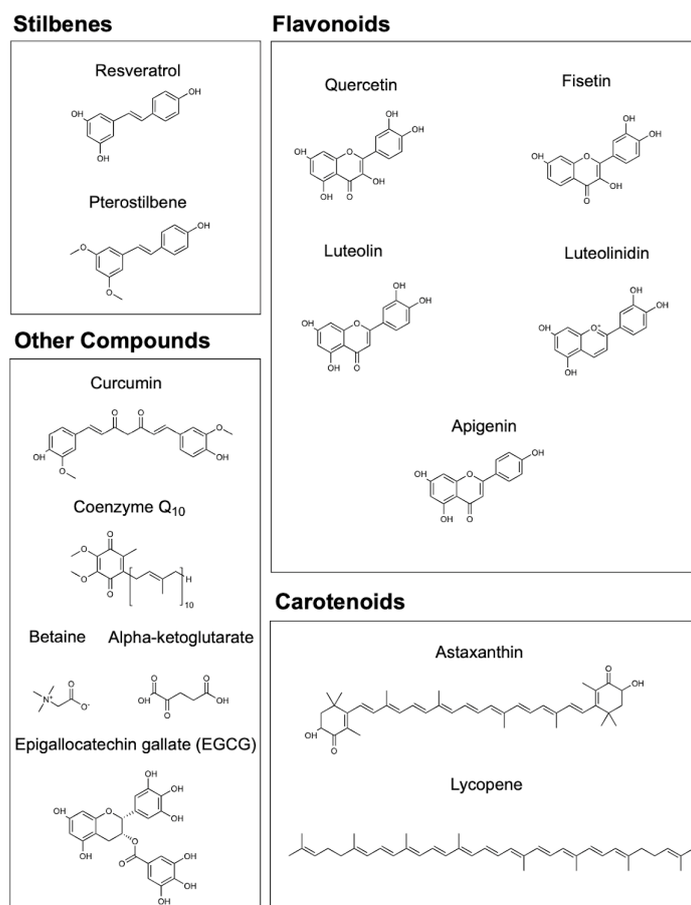


Figure 2. Chemical structures of the aforementioned nutraceutical compounds.

4. Conclusive Remarks

Having established the benefits of numerous ingredients involved in longevity-related pathways, a clearer picture emerges of the entangled molecular web of the NAD⁺/SIRT1/CD38 axis in the context of healthy aging supplementation. Many ingredients on their own show numerous benefits in the realm of many age-related disorders, such as neurodegeneration, cardiovascular and metabolic disease, diabetes, cancer, frailty, sarcopenia, hearing loss, osteoarthritis and osteoporosis, and ocular disorders [187]. CD38 inhibitors may become a focus for the development or screening of new compounds, which may imply longevity induction *in vivo*, and, ultimately, in humans. CD38-inhibitor 78c is one compound that has yet to enter human interventional trials but has shown marked improvements in longevity and physical performance, and metabolic profile in mice [188].

A beneficial outlook may be to further look into other senolytics that ameliorate numerous hallmarks of aging. Further clarification of SIRT1 activity in relation to downstream effects implicated in longevity extensions shown in experimental models is required, particularly in humans, to determine the maximal effect of innervation through nutraceutical interventions. Testing superiority trials of NMN or NR in combination with other ingredients compared to NMN or NR alone, geroprotectors, senolytics alone, or other NAD⁺ precursor molecules will prove vital to maximizing the effects of each ingredient along with maximal NAD⁺ boosting potential. The potential of generated knowledge from further analyzing these relationships within the context of human aging may also offer insight into other pathways that may be implicated in longevity but may yield mixed results in the context of the fine balance for tumor suppression, senescence, and senolytic activity. Subsequently, avenues to target the pathology of aging and age-

related diseases to increase healthspan may arise upon further exploration of other geroprotectors, particularly within clinical trials.

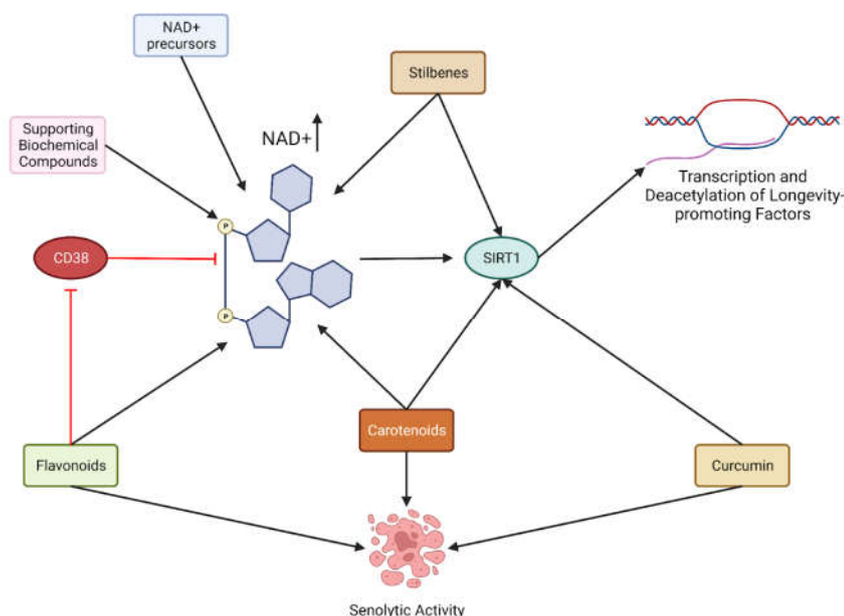


Figure 3. Hypothesized model of supplementing NAD⁺ precursors with other NAD⁺ enhancing geroprotectors. In addition to NAD⁺ precursors to raise NAD⁺ levels and enhance SIRT1 activity, stilbenes are able to support NAD⁺ levels and further activate SIRT1. Many flavonoids retain senolytic and CD38-inhibitory activity and can further innervate NAD⁺ stores. Curcumin and carotenoids retain similar properties in addition to SIRT1 activation. This interacting web of support may result in higher NAD⁺ stores than precursor-alone administration, producing longevity-promoting transcriptional benefits.

By screening for compounds that exhibit an effect on the aging-related molecular pathways, clever formulations of geroprotective ingredients can be formulated, and in particular, produce a synergistic effect of boosting NAD⁺ levels or SIRT1 activity or reducing degrading CD38 activity to maximize lifespan and healthspan extension supplements. Due to the lack of superiority trials analyzing combination therapies and potential molecular synergies, a call for further RCTs is highly warranted in the field of healthy aging orthomolecular medicine and geroprotective nutrition.

Author Contributions: All authors participated in conceptualizing and editing the review. AS and CYE wrote the manuscript in consultation with the other authors.

Funding: This research was funded by the Swiss National Science Foundation: the SNF P3 Project 190072 to CYE.

Acknowledgments: We thank the Ewald Lab and the Maximon Team for feedback and critical reading of the manuscript. Figure 1 and 3 were made with Biorender (Publication licensing numbers: BH24LJRAQG and VB24LJR3ZB). Figure 2 was rendered using Chemdraw, and compounds were sourced from PubChem (<https://pubchem.ncbi.nlm.nih.gov/>).

Conflicts of Interest: AS is a Research and Development Intern at AVEA Life AG. SC is a Co-founder and Chief Product Officer at AVEA Life AG. EMR is a Chief Scientific/Medical Officer and Partner at Maximon AG. CYE is a Co-founder and shareholder of Avea Life AG, and is on the Scientific Advisory Board of Maximon AG, Biotein, Longaevus Technologies LTD, and Galyan Bio, INC.

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