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1	The role of inorganic nitrate and nitrite in cardiovascular disease
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3	Jacklyn Jackson ¹ , Amanda Patterson ² , Lesley MacDonald-Wicks ² and Mark McEvoy ³
4	
5 6	¹ School of Health Sciences, Faculty of Health and Medicine, University of Newcastle, University Drive, Callaghan, NSW, Australia
7 8	² Priority Research Centre in Physical Activity and Nutrition, University of Newcastle, University Drive, Callaghan, NSW, Australia
9 10	³ Centre for Clinical Epidemiology and Biostatistics, Hunter Medical Research Institute, University of Newcastle, Callaghan, NSW, 2309, Australia
11	
12	Corresponding author:
13	Mark McEvoy
14	Centre for Clinical Epidemiology and Biostatistics, Hunter Medical Research Institute,
15	University of Newcastle
16	University Drive, Callaghan, NSW, Australia.
17	E-mail: Mark.Mcevoy@newcastle.edu.au
18	Tel: +61 2 4042 0518
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27 Abstract

Cardiovascular disease is the leading cause of death worldwide, a consequence of mostly poor 28 lifestyle and dietary behaviours. Although whole fruit and vegetable consumption has been 29 consistently shown to reduce cardiovascular disease risk, the exact protective constituents of these 30 31 foods are yet to be clearly identified. A recent and biologically plausible hypothesis supporting the cardio-protective effects of vegetables has been linked to their inorganic nitrate content. 32 Approximately 60-80% inorganic nitrate exposure in the human diet is contributed from vegetable 33 consumption. Although inorganic nitrate is a relatively stable molecule, under specific conditions it 34 can be metabolised in the body to produce nitric oxide via the newly discovered nitrate-nitrite-nitric 35 oxide pathway. Nitric oxide is a major signalling molecule in the human body, and has a key role in 36 37 maintaining vascular tone, smooth muscle cell proliferation, platelet activity and inflammation. Currently, there is accumulating evidence demonstrating that inorganic nitrate can lead to lower 38 39 blood pressure and improved vascular compliance in humans. The aim of this review is to present an informative, balanced and critical review of the current evidence investigating the role of 40 inorganic nitrate and nitrite in the development, prevention and/or treatment of cardiovascular 41 disease. Although there is evidence supporting short term inorganic nitrate intakes for reduced 42 blood pressure, there is a severe lack of research examining the role of long-term nitrate intakes in 43 the treatment and/or prevention of hard cardiovascular disease outcomes, such as myocardial 44 infarction and cardiovascular mortality. Epidemiological evidence is needed in this field to justify 45 continued research efforts. 46

47

48 Introduction

49 Despite major medical research advancements over the past 50 years, cardiovascular disease (CVD) 50 remains the leading cause of death worldwide and is responsible for 39% of non-communicable 51 disease (NCD) deaths in populations aged under 70 years old⁽¹⁾. The leading NCD risk factor is 52 hypertension, which is responsible for 13% of global deaths each year and is a major risk factor for 53 coronary artery disease (CAD), ischemic heart disease (IHD) and stroke⁽¹⁾.

54

The pathogenesis of CVD is influenced by a variety of risk factors that can be broadly categorised as either modifiable or non-modifiable⁽²⁾. Non-modifiable risk factors cannot be controlled through intervention and include advancing age, gender (men at greater risk than pre-menopausal women; post-menopausal women at greater risk than men), ethnicity and family history of CVD⁽²⁾. Modifiable risk factors on the other hand, have the ability to be manipulated through intervention in

60 order to control, treat or modify the risk factor⁽²⁾. Established modifiable risk factors for CVD

include hypertension, tobacco use, raised blood glucose, physical inactivity, unhealthy diet, raised
blood cholesterol/lipids and overweight and obesity⁽²⁾.

Implementation of various lifestyle strategies which target specific modifiable risk factors can
reduce the risk of CVD by up to 80%^(1; 2). Thus indicating that CVD is a chronic and mostly
lifestyle induced disease, to which the majority of current mortality is the consequence of previous
exposures to behavioural risk factors such as inappropriate nutrition, insufficient physical activity
and tobacco exposure ^(2; 3; 4; 5). In addition, excess weight and central obesity, increased blood
pressure, dyslipidaemia, diabetes and low cardiorespiratory fitness are among the factors
contributing principally to CVD risk^(2; 6).

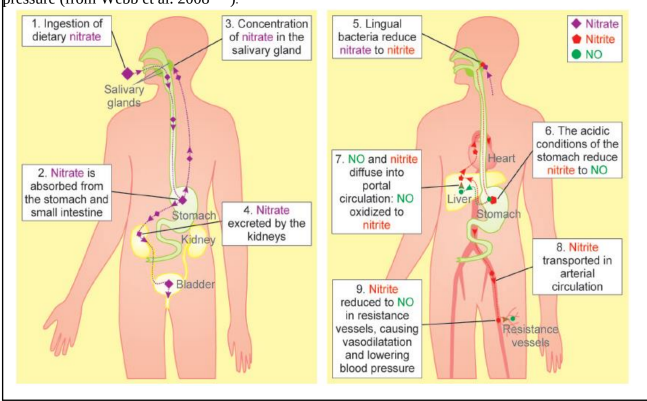
70 Given the scope and prevalence of CVD within our current food and lifestyle environment, it is clear that preventative measures are the most appropriate to deal with this global health issue in 71 order to reduce the costs to both the community (through improved quality of life) and governments 72 through a reduction in hospitalizations, medication use and rehabilitation⁽²⁾. Although behavioural 73 74 factors such as smoking cessation and increased physical activity appear relatively straight forward targets for public health preventative interventions, the definition of a perceived "healthy" diet has 75 76 changed over time leading to a general sense of public confusion and uncertainty surrounding the topic (7; 8). 77

Currently, the most compelling dietary evidence for CVD prevention is linked to whole diet
 approaches such as the Mediterranean and DASH (Dietary Approaches to Stop Hypertension) diets
 ^(7; 9). Although the cardio-protective effects of these diets may be credited to a whole diet/whole
 food effect, some individual nutritive components of these foods have also been extensively
 investigated.

83 The investigation of single nutritive components demonstrates the evidence is less clear, this is especially noticeable for fruit and vegetable constituents. While whole fruit and vegetable 84 consumption has been consistently shown to reduce CVD risk, as evidenced by various prospective 85 studies showing a direct inverse association between fruit and vegetable intakes and the 86 development of CVD events such as myocardial infarction (MI) and stroke^(10; 11; 12; 13), the various 87 constituents of fruits and vegetables such as vitamin C, polyphenols, fibre and antioxidants are yet 88 to clearly demonstrate a beneficial link or a physiological pathway for their individual effect^{(14; 15; 16;} 89 17; 18) 90

A recent and biologically plausible hypothesis for the cardio-protective and blood pressure lowering
effect of vegetables has been linked to their inorganic nitrate (NO₃⁻)/nitrite (NO₂⁻) content ⁽¹⁹⁾.
Support for this hypothesis has been implied in studies indicating that nitrate rich green leafy

- 94 vegetables and vitamin C rich fruits and vegetables contribute most to the apparent cardiovascular
- 95 protective effect of total fruit and vegetable intake ^(20; 21). Additionally, cardio-protective diets
- 96 including the DASH, Mediterranean and Traditional Japanese diets have been shown to naturally
- 97 contain high quantities of inorganic nitrate (147-1222 mg/d) relative to a typical Western style diet
- 98 (~75mg/d) $^{(22; 23; 24)}$.
- 99 Within the human body, inorganic nitrate/nitrite (NO_x) can be metabolised to produce nitric oxide
- (NO) (Figure 1.) ^(25; 26). NO is a highly valuable signalling molecule and has been demonstrated to
- 101 mediate favourable effects on blood pressure control, platelet function, vascular health and exercise
- 102 performance (27; 28; 29; 30). In addition, the utility of inorganic NO_x as a NO donor may be of
- 103 particular relevance given that one serving of nitrate rich vegetables (such as beetroot) has been
- 104 estimated to produce more NO under specific conditions than can be endogenously formed by the
- 105 classical L-arginine-Nitric-Oxide-Synthases pathway each day^(19; 31; 32).
- 106 Currently, the true effect dietary/inorganic NO_x may have on CVD risk factors and outcomes is
- 107 poorly understood, but it is a highly worthwhile line of investigation given that an increased daily
- 108 consumption of nitrate intake represents a potential low cost and simple treatment option for
- 109 reducing CVD burden.
- **Figure 1. The fate of dietary nitrate.** Nitrate is systematically absorbed becoming concentrated in
- the salivary glands and part of the salivary circulation. Salivary nitrate is reduced to nitrite by oral
- bacteria. In the stomach nitrite may produce NO. Nitrite transported in arterial circulation can be
- reduced to NO in low oxygen concentrations which can lead to vasodilation and reductions in blood pressure (from Webb et al. 2008⁽³³⁾).



- 116 **Production of nitric oxide in the body**
- 117 Endogenous production via the L-arginine nitric oxide synthase pathway
- 118 The notion that NO_x could be produced endogenously in the body was first considered in the early
- 119 1980s, upon finding that NO_x excretion was exceeding quantities of ingestion in animal and human
- 120 models ^(34; 35). Later it was demonstrated that L-arginine was the substrate for synthesizing nitrogen
- 121 oxides endogenously via the action of NO synthase (NOS) enzymes ⁽³⁶⁾.
- 122

In healthy individuals the L-arginine-NOS pathway can produce sufficient quantities of NO to
maintain health (approximately 1.7 mmol/day)^(31; 32). However, conditions such as diabetes mellitus,
aging, hypercholesterolemia and tobacco exposure have been found to impact the bioactivity of
endogenously produced NO via one or more of the following functions ^(37; 38; 39; 40; 41; 42):

- Increased degradation of NO^(38; 42; 43)
- Altered phosphorylation and activation of NOS ^(38; 43)
- Increased production of NOS inhibitors (eg. Asymmetric Dimethylarginine (ADMA)),
 leading to disruption of NOS activation ^(38; 39; 41; 42; 43)
- Deficiency of NOS substrate, L-arginine ^(34; 38; 41)
- Reduced availability of one or more cofactors essential for NOS function ^(34; 38)

While appropriate medical management, consumption of a healthy diet and moderate exercise can 133 somewhat reverse these effects, it has been postulated that supplementing portions of the NOS 134 pathway may enhance NOS activity and NO production ^(38; 41; 43). This has been of particular 135 importance given that increased ADMA levels inhibit NOS function and has been cited as the 136 strongest risk predictor of cardiovascular events, and all cause and cardiovascular mortality in 137 people with CAD⁽⁴⁴⁾. Although it remains unclear whether a change in ADMA can alter CVD risk, 138 interventions such as L-arginine supplementation have been shown to improve endothelial-mediated 139 vasodilation in people with elevated ADMA levels^(41; 44). 140

- 141 As a result, the effect of L-arginine supplementation has been investigated and short term
- supplementation was shown to improve endothelial function and relieve symptoms in patients with
- 143 coronary heart disease⁽⁴⁵⁾. Long-term (6 months) supplementation however, demonstrated no
- beneficial effect⁽⁴⁶⁾. In fact the long-term L-arginine supplementation lead to increased rates of
- 145 death and less cardiovascular improvements compared to the placebo due to the development of
- arginine toxicity and hyperkalemia (abnormally high serum potassium)^(47; 48). In addition, the utility
- 147 of supplementing arginine is questionable given that arginine is classified as a "semi essential" or
- 148 "conditionally essential" amino acid, depending on the developmental stage or health status of the

individual⁽⁴⁹⁾. However, it is generally accepted that healthy adults should not need to supplement
with arginine as their bodies produce physiologically sufficient amounts⁽⁴⁸⁾. Arginine is also highly
abundant in the diet, as rich dietary sources include meat, dairy, vegetables, legumes and
wholegrains^(48; 49).

The "arginine paradox" appears to address this notion, as it refers to the phenomenon that 153 exogenous arginine causes NO mediated biological effects, despite the fact that NOS are 154 theoretically saturated in the substrate L-arginine ⁽⁴⁹⁾. A recently published cross-sectional study 155 including 2771 men and women investigated whether regular dietary intakes of L-arginine were 156 associated with serum NOx, as an indicator of systemic NO production ⁽⁵⁰⁾. This study found that 157 increased dietary L-arginine intakes were strongly associated with serum NOx, which was 158 independent of the overall dietary patterns of the study participants and other dietary factors, 159 including intakes of high nitrate containing foods (likely due to collection of fasting blood samples) 160 ⁽⁵⁰⁾. Therefore, although there may be some utility in consuming adequate amounts of arginine, 161 which is readily achieved by consumption of a healthy balanced diet, there appears to be no great 162 benefit for the general population to be using arginine supplements. However, dietary intervention 163 to also consume nitrate rich foods holds much promise for supplementing the NOS pathway via the 164 alternative nitrate-nitrite-NO pathway. 165

166 **The nitrate-nitrite-NO pathway:**

167 Up until the early 1990s, plasma NO_x were considered to be biologically inactive end products of 168 NO production in the human body. However it is now clear that under specific conditions nitrate 169 and nitrite anions can be recycled in vivo back to $NO^{(26; 27; 51; 52)}$.

170

With a bioavailability of 100%, ingested inorganic nitrate is swiftly absorbed in the proximal small 171 intestine leading to significantly raised plasma nitrate concentrations for a period of up to 5-6 hours 172 post nitrate ingestion (27; 33; 53; 54; 55). About 75% of this nitrate is excreted at the kidneys, however 173 the other 25% of plasma nitrate is actively extracted by the salivary glands, leading to salivary 174 nitrate concentrations which are 10-20 times higher than plasma nitrate concentrations (27; 43; 55; 56; 175 ⁵⁷⁾. Salivary nitrate accumulation must occur in order for nitrate to be reduced to nitrite, as 176 anaerobic bacteria in the oral cavity use nitrate as an alternative electron acceptor to oxygen during 177 respiration^(27; 55; 56; 58). When this nitrite rich saliva is swallowed it is reduced in the acidic stomach 178 to produce nitrogen oxides including NO^(26; 27; 52; 59). Today, this process is widely known as the 179 nitrate-nitrite-NO pathway, and is thought to be one of the body's major sources of NO generation, 180 181 especially in situations when NO bioavailability via the conventional L-arginine-NOS pathway is compromised. In addition it has been suggested that the nitrate-nitrite-NO pathway may play a 182

- significant role in maintaining levels of bioactive NO and may be critical for maintaining
- 184 cardiovascular homeostasis in the body $^{(27; 53; 60)}$.
- 185 Noteworthy factors other than inorganic nitrate and nitrite consumption which have been shown to186 facilitate the nitrate-nitrite-NO pathway include:
- The entero-salivary nitrate cycling: Approximately 25% of plasma nitrate is actively
 taken up by the salivary glands leading to significant nitrate accumulation in the saliva.
 Within the oral cavity, anaerobic bacteria reduce nitrate to nitrite via the action of nitrate
 reductive enzymes. Nitrite rich saliva must be swallowed to produce NO in the acidic
 stomach.
- The importance of this salivary nitrate cycling has been demonstrated in studies where subjects spat after a dietary load of inorganic nitrate, preventing the opportunity for nitrate to accumulate in the saliva and be reduced to nitrite, therefore preventing NO production and any beneficial effects ^(25; 33; 61).
- Presence of anaerobic bacteria: Mammalian bacteria can utilise nitrate as an alternative 196 • 197 electron acceptor to oxygen during respiration, and is a vital component of the nitrate-nitrite-NO pathway as human cells lack the required nitrate reductase enzymes ⁽⁶¹⁾. The importance 198 199 of these bacteria has been further established in studies of germ free rats, in which gastric NO formation was negligible post dietary nitrate load ⁽⁶²⁾. Additionally, human studies have 200 demonstrated that the use of commercial antibacterial mouthwash in humans abolished any 201 202 blood pressure lowering effects of a dietary nitrate load indicating that the mouthwash killed off the commensal facultative bacteria in the mouth, thus preventing the production of nitrite 203 and NO leading to a loss of beneficial health effects (63; 64; 65). 204
- Hypoxic conditions: The rate in which nitrate is reduced to nitrite is 30 times greater during conditions of low oxygen tension, as the oral bacteria use salivary nitrate as an alternative electron acceptor to oxygen during respiration⁽⁶⁵⁾. Xanthine oxidoreductase (XOR) has also been shown to catalyse the reduction of nitrite to NO in hypoxic conditions ^(66; 67; 68).
 This could also account for the increased production and utility of NO seen in exercising skeletal muscle or during myocardial ischemia ^(52; 61; 69).
- 211 It is also important to note that plasma nitrite can be reduced to NO along the physiological 212 oxygen gradient of the circulatory system⁽⁷⁰⁾. Specifically, deoxygenated haemoglobin in
- the peripheral circulation can act as a nitrite reductase for NO production, as it has been
- revealed that as haemoglobin deoxygenation increases, more NO is produced ^(71; 72; 73). This
- provides an explanation for how various human studies have observed vasodilation post a
- 216 NO_x load, in healthy subjects at rest $^{(33;74)}$.

- Acidic conditions: Nitrite in the acidic stomach has been shown to spontaneously
 decompose to NO, a reaction that appears to increase in conditions of reduced pH (increased acidity)⁽²⁶⁾. The importance of an acidic stomach for this reaction has been demonstrated in a study, showing that NO production via nitrite protonation was inhibited in individuals
 using proton pump inhibitors (medications which reduce the acidity of gastric juices)⁽⁷⁵⁾.
- Presence of reducing agents including vitamin C and polyphenols: Both vitamin C and polyphenols are abundant in a vegetable rich diet, and their presence in the diet has been shown to favour the formation of NO via the nitrate-nitrite-NO pathway and prolong the half-life of NO in the stomach ^(76; 77).
- 226

227 Sources of dietary inorganic nitrate and nitrite:

Nitrogen is vital to life on Earth and can undergo many chemical and biological changes in order to
 be amalgamated into living and non-living material. An essential form of environmental nitrogen
 includes inorganic nitrate, as an adequate nitrate supply in the soil is essential for plant growth ^(43; 78).

The two major determining factors of the nitrate content of vegetables and fruit, include their 232 species and the amount of available nitrate in the soil ⁽⁴³⁾. Some species of vegetables such as green 233 leafy vegetables (mean nitrate ~ 975-3624 mg/kg) and beetroot (mean nitrate ~ 1992 mg/kg) are 234 naturally high in nitrate, however environmental factors can lead to great variation among samples 235 ⁽²²⁾. These factors include seasonal differences and disruption to normal plant growth, leading to 236 nitrate accumulation in the plant leaves, stems and stalks, due to changes in the photosynthetic 237 conversion of plant nitrate to amino acids ^(78; 79; 80). Therefore, established factors shown to effect 238 the normal growth of plants include drought conditions, high temperatures, shady and cloudy 239 conditions, deficiency of soil nutrients, and excessive soil nitrogen ⁽⁴³⁾. Additionally, farming 240 practices leading to damaged produce, early harvest, storage and transport conditions, processing 241 and cooking practices will also result in significant variation in vegetable and fruit nitrate content 242 (43) 243

European based studies have demonstrated that organically grown vegetables have a lower nitrate content than conventionally grown crops, despite the fact that organic fertilizers may cause high nitrate levels in vegetables, depending on the types and amount of organic fertilizers applied ⁽⁸¹⁾. A California based study by Muramoto et al (1999) reiterated this notion, as it found spinach grown and harvested during the same season and under the same farming practices had a wide range of nitrate contents. This range appeared greatest in organic spinach, in which the maximum nitrate content measured was 3000 mg/kg, which was five times higher than the minimum (600 mg/kg) ⁽⁸¹⁾. However, this study also demonstrated that conventionally grown spinach contained on average
30% more nitrate than spinach grown organically, a result most likely explained due to the wide use
of nitrogen containing fertilizers in conventional farming ⁽⁸¹⁾.

Muramoto et al. also found a statistically significant seasonal difference in the nitrate content of 254 iceberg lettuce, as winter samples were found to have on average 52% more nitrate than summer 255 samples⁽⁸¹⁾. This finding is consistent with Ekart et al (2013), which found lettuce harvested during 256 summer had a statistically significant lower nitrate content than lettuce harvested during winter 257 (summer harvest: 1209 mg/kg, winter harvest: 2164 mg/kg)⁽⁸²⁾. In addition, Ekart et al found that 258 washing leafy greens reduced the nitrate content of foods on average by 19%. Other processing 259 such as boiling, blanching and sautéing, were found to significantly reduce the nitrate content of 260 spinach by 53%, 36% and 30% respectively⁽⁸²⁾. A finding which could be partly explained due to 261 the water soluble nature of inorganic nitrate⁽⁸³⁾. 262

Due to the high variability of nitrate within plant species, accurate and reliable nitrate intake
measured from fruit and vegetable consumption is difficult to predict. Despite this, combined
vegetable and fruit intake is the major source of exogenous inorganic nitrate exposure and are
predicted to constitute 30-90% of total nitrate intake ⁽⁸⁴⁾. Other sources of nitrate intake include
drinking water and meat products, however their nitrate content is highly regulated to comply with
strict government limits ^(85; 86; 87; 88; 89).

Nitrate occurs naturally in the water supply, however in most developed countries water nitrate is generally present in concentrations much lower than allowed in the water guidelines ($\leq 50 \text{ mg/L}$) ^(85; 86; 88). Therefore, nitrate from the water supply is unlikely to contribute significantly to total nitrate intake in comparison to food sources.

Nitrate and nitrite salts (e.g. potassium nitrite/ sodium nitrate) have been used as food additives in cured meats for many years due to its effectiveness in ensuring microbial safety and its ability to enhance the flavour and appearance of the product ⁽⁴³⁾. The maximum levels of nitrate and nitrite allowed as a food additive have been defined (Table 1) ^(85; 90; 91; 92).

It has been estimated that approximately 60-80% of dietary nitrates are derived from vegetables (mainly green leafy and root vegetables) indicating that vegetable intake tends to contribute the greatest quantities of dietary nitrate (Table 2) ^(22; 93). This has been further implied by dietary patterns such as the DASH diet, Mediterranean, vegetarian and traditional Japanese diets which tend to include high quantities of vegetables (5 or more serves/d) and provide approximately 147-1222 mg nitrate per day ^(22; 23; 24). This is a relatively high nitrate intake compared with the typical Western style diets which tends to be low in vegetables (1-3 serves/d) and provides around 60-75 mg nitrate per day ⁽²⁴⁾. In addition, processed and cured meats are frequently cited as the major dietary source of nitrite (Table 3) ^(22; 25; 84; 94), followed by various fruits and vegetables (Table 2, 4 and 5), which have been physically damaged or poorly stored as enzymes present in the plant tissues and/or contaminating bacteria facilitate the reduction of nitrate to nitrite ^(43; 85).

288

289 Nitric oxide in the cardiovascular system:

Within the cardiovascular system, basal endothelial NO has a critical role in maintaining 290 cardiovascular health as it controls vascular tone, smooth muscle cell proliferation and growth, 291 platelet activity and aggregation, leukocyte trafficking, expression of adhesion molecules, and 292 inflammation ^(34; 94; 95; 96; 97; 98; 99). However, when the bioavailability of NO is compromised, the 293 beneficial effects of NO are lost and endothelial dysfunction predominates due to the imbalance 294 created between the release of vasoconstrictors and vasodilators (such as NO)^(53; 100; 101). This idea 295 has been supported in a study conducted by Kleinbongard et al (2005) which found that plasma 296 297 nitrite levels are a reliable indicator of endothelial dysfunction and correlate with cardiovascular risk factors in humans⁽¹⁰²⁾. Additionally, endothelial dysfunction has been strongly linked with 298 atherosclerosis development and a number of cardiovascular disorders such as hypertension, 299 coronary artery disease, congestive heart failure and peripheral artery disease in multiple 300 longitudinal studies (53; 101; 103; 104; 105; 106; 107). 301

While in the past most of the evidence suggesting a relationship between endothelial dysfunction 302 and clinical events from atherosclerosis development were considered "circumstantial", more 303 recently conducted cross-sectional studies have indicated that severe endothelial dysfunction of the 304 arteries can trigger events of unstable angina and myocardial infarction^(108; 109). Al Suwaidi et al 305 (2000) studied 157 patients with mild coronary artery disease for 2.3 years, and found an increased 306 incidence of cardiovascular events in patients with impaired endothelium-dependent vasodilation 307 (NO production of endothelium) of the coronary arteries ⁽¹⁰⁴⁾. In another study by Katz et al (2005), 308 259 subjects with chronic heart failure were assessed prospectively, to which endothelial 309 dysfunction in chronic heart failure was found to significantly increase risk of mortality (110). Thus 310 supporting the notion that coronary endothelial dysfunction plays a role in the pathogenesis of 311 coronary atherosclerosis, risk of cardiac events and death (104; 110). 312

313 Many factors are known to predispose endothelial dysfunction, due to reductions in NO

314 concentrations and bioavailability in humans ^(34; 111; 112). These factors are consistent with the

modifiable and non-modifiable risk factors for CVD, including hypertension, hypercholesterolemia,

diabetes, tobacco use, physical inactivity, consumption of unhealthy diets and increased age and

- 317 gender (NO bioavailability is reduced in post-menopausal women, a period in which CVD risk is
- drastically increased in women)^(34; 112; 113; 114; 115; 116; 117; 118; 119; 120). Interestingly, improved
- endothelial function is a common feature of experimental intervention studies, which have been
- 320 shown to reduce cardiovascular risk and improve endothelial dependent vasodilation in the coronary
- and peripheral circulation ⁽¹⁰⁸⁾. Such interventions commonly include use of lipid and blood pressure
- lowering medications, smoking cessation and increased physical activity^(108; 117; 121; 122; 123; 124).
- 323 However, the notion that inorganic nitrate and nitrite either consumed from dietary sources such as
- 324 green leafy vegetables or supplement is relatively new, and their therapeutic potential as a NO
- donor via the nitrate-nitrite-NO pathway remains unclear^(112; 125).

326

327 Cardiovascular protective actions of nitric oxide:

Nitric oxide is non-polar and can diffuse freely across cell plasma membranes and is a key
signalling molecule capable of many important functions acting primarily by stimulating intra-

330 cellular receptors within the target cell $^{(126)}$.

Within the vasculature of the cardiovascular system, the primary role for NO's action is for the regulation of vascular function and blood pressure, a notion which has been clearly demonstrated in animal models in which synthesis of NO was blocked leading to persistently elevated blood pressure ^(112; 127). In addition, this interaction has been demonstrated in some recently conducted short-term dietary nitrate trials in humans, which showed that peak blood pressure lowering effects were achieved in synchronization with peak plasma concentrations of nitric oxides (NOx) post a dietary nitrate load ^(28; 33; 128).

- The cellular pathway in which NO exerts this vasodilatory action is well established. Nitric oxide rapidly diffuses across vascular smooth muscle cell membranes. Within the smooth muscle cells, NO binds to and activates guanylyl cyclase to produce cyclic guanosine monophosphate (cGMP) (¹²⁶⁾. Once produced, cGMP can have a number of effects in the cells, but many of these effects are mediated thought the activation of protein kinase G (PKG). Activation of PKG via cGMP leads to the activation of myosin phosphatase which in turn leads to smooth muscle cell relaxation and vasodilation (^{126; 127)}.
- 345 In addition to regulating vascular tone, NO can facilitate many other important functions preventing
- the development of atherosclerosis, which include antiplatelet effects, anti-proliferative effects,
- anti-inflammatory, and anti-oxidant effects ^(127; 129; 130). Although the cellular pathways for these
- 348 actions are yet to be clearly defined, it is clear that NO is capable of binding to or reacting with a

349 variety of chemical modalities within the cellular environment, including metal containing proteins,

350 membrane receptors, ion channels, enzymes, transcription factors and oxygen species ^(127; 131).

351

352 Other Nitric Oxides and Possible Mechanisms in the Cardiovascular System:

While NO is the most widely cited bioactive metabolite underpinning the cardiovascular therapeutic benefits of dietary inorganic nitrates and nitrites, it has been suggested that other nitric oxides also play a role ^(25; 93). This may be expected, given that dietary constituents in the stomach may react with each other in order to form a variety of bioactive compounds ⁽²⁵⁾. Examples of such compounds include; nitrated fatty acids, nitrosothiols and ethyl nitrite ⁽²⁵⁾.

358

359 While the biological significance of these compounds are yet to be made clear, the following 360 actions have been suggested:

- Ethyl Nitrite: Rat models have shown that ethanol from alcoholic drinks can interact with
 salivary-derived nitrite in the acidic stomach leading to the production of ethyl-nitrite ^(25; 132).
 Ethyl-nitrate is a potent smooth muscle relaxant and may have a vasodilatory role in the
 cardiovascular system ⁽¹³²⁾.
- Nitrosothiols: In the stomach, nitrite has been shown to induce S-nitrosation within the gastric compartment. S-nitrosothiols are thought to represent a circulating endogenous
 reservoir of NO acting as a NO donor ⁽²⁵⁾.
- Nitrated Fatty Acids (nitroalkenes): Nitrogen oxides can react with unsaturated fatty acids to produce nitroalkenes. Analysis of synthetic nitroalkenes derivatives of oleic, linoleic and arachidonic acid reveals that these species possess unique chemical reactions which may support multiple cell signalling events such as vasodilation and reduced inflammation ⁽²⁵⁾.
 Such events may be mediated through their NO donor capabilities.
- 373

Currently the systemic capabilities of these bioactive nitrogen compounds remain uncertain, however it highlights a possible whole diet effect for exerting a beneficial effect on NO and other relevant cardiovascular signalling molecules. This notion is highlighted by Lundberg and Weitzberg (2010), indicating that various dietary constituents of the Mediterranean diet may interact in the stomach to produce these potentially therapeutic compounds, and may provide an additional explanation for the cardiovascular health benefits/protection seen with this dietary pattern ^(25; 93).

- 380
- 381

382 Inorganic versus organic nitrate and nitrite

Organic nitrates such as glyceryl trinitrate (GTN) and isosorbide mononitrate represent the first class of NO donors to reach the clinical setting and have been used extensively in the treatment of various cardiovascular conditions including angina, coronary artery disease and heart failure ⁽⁸³⁾.

Unlike inorganic nitrates which are relatively simple molecules and naturally occurring in fruits and vegetables, organic nitrates are synthetic compounds produced by a reaction between nitric acid and an alcohol group ⁽⁸³⁾. Organic nitrates are complex, non-polar hydrocarbon chains attached to a nitrooxy-radical (-ONO₂), which is responsible for its biological effects (Table 6.) ⁽⁸³⁾.

Once organic nitrates are introduced to the blood system, levels rise quickly leading to the rapid

onset of their action $^{(83)}$. At low doses (≤ 1.25 mg/kg body weight) organic nitrate has been

demonstrated to dilate large conductance veins and large arteries. While at high doses (2.5-5mg/kg

body weight) organic nitrates can also induce dilation of the arterioles of the microcirculation $^{(83)}$.

395 These vasodilatory effects of organic nitrates have been shown to reduce cardiac work and lower

myocardial oxygen requirements, which may alleviate or even prevent cases of myocardial

- infarction $^{(133)}$. In addition, it has been suggested that organic nitrates have anti-aggregatory
- 398 properties in patients with stable and unstable angina $^{(133)}$.

399 Today in clinical practice short acting organic nitrates most notably in the form of GTN are administered during the symptomatic treatment of myocardial infarction and angina^(83; 133). Glyceryl 400 tri-nitrates are generally administered in the form of either a mouth spray or intravenous infusion, to 401 which onset of action is rapid (2-3 minutes)⁽¹³³⁾. Although short term treatment with organic 402 nitrates has some positive impact on endothelial function, acute side-effects of their use include 403 hypotension, dizziness, nausea and headache⁽⁸³⁾. Also, despite the high potency of organic nitrates 404 and their long history as being used to treat various cardiovascular diseases, nitrate tolerance is a 405 huge limitation and an undesirable side effect of their use (83; 133). 406

407 Nitrate tolerance is a complex phenomenon and is poorly understood, however it is clearly a result
408 of chronic organic nitrate use to which nitrovasodilator-responsiveness is lost ⁽⁸³⁾. Nitrate tolerance
409 has been reported to occur within 1-3 days of continuous GTN treatment in patients with

410 myocardial infarction, stable angina and chronic congestive heart failure ⁽¹³³⁾. Further, chronic

411 organic nitrate use has also been linked to endothelial dysfunction, increased production of free

radicals and development of vascular tolerance to other endothelium dependent vasodilators ⁽⁸³⁾.

Although this phenomenon is poorly understood, recent animal and human studies indicate that

414 increased vascular production of the superoxide anion (O_2^-) underlies the mechanism for tolerance

415 ⁽¹³³⁾. This oxidative stress hypothesis of nitrate tolerance is supported by numerous reports

- 416 demonstrating that the tolerance is prevented by co-administration of antioxidants (eg. vitamin C,
- 417 vitamin E and folic acid) and interventions which inhibit reactive oxygen species (ROS) formation
- 418 (lipid and blood pressure lowering medications)^(133; 134; 135; 136).
- 419 It is interesting to note that the phenomenon of tolerance is not exhibited with the consumption of
- 420 inorganic nitrates/nitrites, however despite showing promise in preventing or treating certain
- 421 cardiovascular conditions, such as hypertension, they have received little attention by the medical
- 422 community $^{(27)}$.

423 Inorganic nitrate and nitrite: From dietary contaminant to potential therapeutic nutrient

- 424 Throughout history, cases of accidental toxic exposure to nitrate and nitrite have been documented,
- 425 however the health risk of excessive inorganic nitrate and nitrite consumption appears specific to
- 426 population subgroups ⁽²²⁾. One of these subgroups includes infants aged less than 6 months, to
- 427 which excessive nitrite exposure has been linked to cases of methemoglobinaemia (blue baby
- 428 syndrome)⁽¹³⁷⁾. As a result, strict regulatory limits have been established to govern the nitrate/nitrite
- content of the drinking water supply and their use as an additive to processed and cured meats in
- 430 order to limit exposure to the population $^{(85; 86)}$.
- 431 Methemoglobinaemia can occur when nitrite oxidises ferrous iron (Fe^{2+}) in haemoglobin to the
- 432 ferric state (Fe^{3+}), resulting in methemoglobin. Methemoglobin is incapable of binding molecular
- 433 oxygen, and impairs oxygen delivery to the tissues causing hypoxia and cyanosis $^{(137)}$. While most
- cases of methomeoglobinaemia have been attributed to the consumption of well water (prone to
- high nitrate accumulation) used for the preparation of infant formula, there have been reported cases
- 436 of nitrate poisoning in infants from the ingestion of plant nitrates ^(86; 137). While Martinez et al found
- that the use of certain high nitrate vegetables (herbs and green leafy vegetables) in infant homemade
- 438 vegetable pureé increased methemoglobinemia in infants (herbs: OR 5.2; 95% CI 1.1-24.6; and
- 439 green leafy vegetables: OR 2.0; 95% CI 0.4-8.7), the most important factor increasing
- 440 methemoglobinemia was the time lapse between vegetable pureé preparation and consumption (OR
- 441 17.4, 95% CI 3.5-86.3 if pureé was prepared 24-48 hrs before and OR 24.9; 95% CI 3.3-187.6 if
- 442 prepared >48 hours before) $^{(138)}$.
- 443 To date human nitrate and nitrite exposure studies have failed to prove a direct link with
- 444 methemoglobinaemia, suggesting that nitrate/nitrite exposure alone may not be responsible for
- 445 methemoglobinaemia development ^(139; 140).
- Another population subgroup that is thought to be at health risk due to excessive nitrate/nitrite
- 447 exposure are high consumers of cured and processed meats ^(22; 141). It has been theorised that nitrates
- 448 and nitrites from processed meats generate N-nitroso compounds which can be carcinogenic $^{(142)}$.

In October 2015 the International Agency for Research on Cancer (IARC) summarized more than 449 800 studies conducted globally, and determined that 50 grams of processed meat each day increased 450 the risk of colorectal cancer by 18%, and therefore concluded that processed meats are carcinogenic 451 ⁽¹⁴¹⁾. In animal studies N-nitrosamines and related N-nitrosamides have been shown to be 452 carcinogenic in a variety of molecular structures ^(143; 144). However, such direct evidence 453 demonstrating nitrate and nitrite as human carcinogens is severely lacking. This has been reflected 454 in the conclusions of the Food and Agriculture Organisation expert committee who found no 455 consistent increased risk of cancer with increasing consumption of nitrate, as available 456 epidemiological studies did not provide evidence that nitrate is carcinogenic to humans⁽¹⁴⁵⁾. 457 Currently, researchers are interested in understanding whether the health risks associated with 458 inorganic nitrates/nitrites outweigh the recently discovered health benefits, however there is a 459 growing consensus that any weak and inconclusive data on inorganic nitrate/nitrite and cancer 460 associations are far outweighed by the potential health benefits of restoring NO homeostasis (22; 84; 461 ^{139; 143)}. In particular this has been demonstrated in various animal and human experimental studies, 462 in which inorganic NO_x has been shown to improve outcomes such as blood pressure, endothelial 463 function, platelet function, ischemia reperfusion injury, exercise performance and host defence (143; 464

465 ^{146; 147; 148; 149; 150; 151}.

466 Evidence of cardiovascular benefit from animal studies:

Intakes of dietary inorganic nitrate have been shown to be strongly cardio protective in animal 467 studies. Carlström et al (2011) indicated this in a four arm dietary intervention trial in rats ⁽¹⁵²⁾. The 468 rats were placed on either a normal salt diet (control); a high salt diet; a high salt diet supplemented 469 with a nutritional (low) dose of nitrate; and a high salt diet supplemented with a pharmacological 470 (high) dose of nitrate for 8-11 weeks⁽¹⁵²⁾. As expected, results demonstrated that chronic 471 consumption of a high salt diet develops hypertension, however when combined with a low nitrate 472 dose, blood pressure was non-statistically significantly lower⁽¹⁵²⁾. On the other hand, the higher 473 nitrate dose lowered blood pressure by a significant 24 mm Hg compared with the plain high salt 474 diet, a magnitude of blood pressure reduction considerably magnified compared with blood pressure 475 reductions observed in another study of healthy normotensive rats using the same nitrate dose (152; 476 ¹⁵³⁾. Similar results were reported by Kanematsu et al. finding that in hypertensive rats, 477 antihypertensive effects were only apparent with the highest dose of nitrate, yet there was a strong 478 tissue protective effect seen with lower doses equivalent to modest dietary intakes ⁽¹⁵⁴⁾. Ferguson et 479 al. demonstrated clinically significant reductions in mean arterial pressure with beetroot juice (BJ) 480 supplementation in exercising rats (Control: 137 ± 3 ; BJ: 127 ± 4 mm Hg, P<0.05), indicating that 481

clinically significant blood pressure reductions may be achievable in doses attained from dietary
 sources ⁽¹⁵⁵⁾.

484

In addition to significant blood pressure control, Carlström el at (2011) found dietary nitrate
supplementation can partly prevent the development of cardiac hypertrophy and high nitrate doses
significantly reduced the fibrotic changes which were observed in the high salt group, two factors
which are major predictors of heart failure ⁽¹⁵²⁾. Two other studies found mice ingesting inorganic
nitrate lead to a significantly reduced infarct size during myocardial ischemia, an important finding
given that reduced infarct size is associated with lower heart failure risk post myocardial infarction
and mortality ^(156; 157; 158).

492 When Baker et al. treated rats with an intra-venous bolus of sodium nitrite across various doses

(0.04, 0.4, 1.0, 4.0, 7.0 and 10.0 mg/kg), prior to initializing a blockage of the coronary artery, there 493 was a clear doses dependent effect of nitrite on infarct size⁽¹⁴⁹⁾. However, it was intriguing to note 494 protection was only found in doses up to 4.0 mg/kg, an effect which was absent at higher doses⁽¹⁴⁹⁾. 495 Rats administered with 4.0 mg/kg nitrite exhibited a significant 32% reduction in infarct size 496 compared to controls (149). Nitrite was also found most effective when administered before and/or 497 during the ischemic event, but not at the onset of reperfusion⁽¹⁴⁹⁾. Further, equivalent doses of 498 sodium nitrate had no effect on infarct size⁽¹⁴⁹⁾. Indicating that administration timing and doses are 499 key considerations for nitrite protection from MI⁽¹⁴⁹⁾. 500

Thrombosis is largely a result of platelet adhesion, activation and aggregation, and is a common
pathology underlying ischemic heart disease and ischemic stroke ^(159; 160). Nitric oxide plays a key
role in preventing thrombosis development⁽¹⁶¹⁾. Park et al. demonstrates this notion upon
discovering an inverse correlation between NO_x levels and platelet activity/aggregation in mice ⁽¹⁶¹⁾.
In addition, Apostoli et al. examined the effect of inorganic nitrite on platelet aggregation in eNOS

deficient mice ⁽¹⁶²⁾. This study found that inorganic nitrite exerts an antiplatelet effect during eNOS
deficiency and suggest that dietary nitrate may reduce platelet hyperactivity during endothelial
dysfunction ⁽¹⁶²⁾.

- Pulmonary hypertension can lead to the remodelling of the artery wall causing abnormalities of
 elastic fibres, intimal fibrosis and medial hypertrophy⁽¹⁶³⁾. This can result in vascular stiffness and is
 a condition linked to the development of chronic heart failure⁽¹⁶³⁾. Sodium nitrite interventions in
 lamb and mice models have shown reductions in pulmonary hypertension specifically during
- 513 hypoxic conditions ^(164; 165). However, Casey et al. found intravenous injections of sodium nitrite
- during normoxic-conditions could lead to reductions of pulmonary and systemic arterial pressure
- and increased cardiac outputs in adult male rats $^{(166)}$. This suggests that sodium nitrite may have a

role in reducing the workload of the heart during pulmonary hypertension and thus protects the
heart and vascular system from associated damage and dysfunction ⁽¹⁶⁶⁾.

Hendgen-Cotta et al. pre-treated mice with nitrate before inducing chronic limb ischemia, and
nitrate supplementation was found to enhance revascularization and increased mobilization of
circulating angiogenic cells (CACs), which are important for the recovery and maintenance of
healthy endothelial function ⁽¹⁶⁷⁾. Heiss et al. on the other hand injected inorganic nitrite into healthy
mice, and found that nitrite significantly increased CACs at 1 hour compared with controls ⁽¹⁶⁸⁾. It is
interesting to note however that when this test was repeated in eNOS deficient mice, no CAC
mobilization was observed, indicating that NOS may be required to take part in nitrate-mediated

525 CAC mobilization ⁽¹⁶⁸⁾.

526 In a study conducted by Sindler et al. the effect of nitrite in aged, but healthy mice was investigated

527 and high dietary nitrite doses were found to reverse age-related vascular dysfunction, arterial

528 stiffness and reduce levels of oxidative stress ⁽¹⁶⁹⁾. This is in line with Carlström et al (2011) which

found key plasma and urinary oxidative stress markers (MDA, $iPF2\alpha$ -VI and 8-OHdG) were

significantly reduced (despite co-consumption of a high salt diet) with both low (0.1 mmol

nitrate/d) and high (1.0 mmol nitrate/d) dose dietary nitrate supplementation, which may be useful

in preventing NO degradation and endothelial dysfunction $^{(152; 170)}$. An interesting finding, given that

oxidative stress is directly linked with an inflammatory response which is thought to have a central
 role in the development of atherosclerosis ⁽⁹³⁾.

Stokes et al. found that mice fed cholesterol-enriched diets for three weeks tend to develop clear 535 signs of vascular disease pathology, including elevated leukocyte adhesion and endothelial 536 dysfunction, an effect which was prevented with nitrite supplementation in the drinking water ⁽¹⁷¹⁾. 537 In another study by Carlström et al (2010) it was demonstrated that several features of metabolic 538 syndrome (including visceral fat and circulating triglycerides, which are strong risk factors for 539 cardiovascular disease) can be reversed by dietary nitrate supplementation, in amounts which 540 correspond to those derived from eNOS under normal healthy conditions or a vegetable rich diet 541 (172) 542

543 Evidence of cardiovascular benefit from human studies:

In 2003 Cosby et al. conducted one of the first studies demonstrating a relationship between inorganic nitrite supplementation and blood pressure reductions in healthy human subjects ⁽⁷¹⁾. This study chose to use sodium nitrite (NaNO₂⁻) infusions providing approximately 75 mg NaNO₂⁻ over two 15 minute periods, a dose which was found to significantly reduce mean blood pressure by 7 mm Hg (P<0.01) ⁽⁷¹⁾. Similar findings were later established using sodium nitrate (NaNO₃⁻) in a

study conducted by Larsen et al. $^{(173)}$. In this study healthy subjects consumed NaNO₃⁻ (8.5) 549 mg/kg/day for 3 days) as a dietary supplement, and although systolic blood pressure was not 550 changed during this time compared with placebo (sodium chloride), diastolic blood pressure was 551 significantly reduced on average by 3.7 mm Hg (P<0.02) and mean arterial pressure was lowered by 552 3.2 mm Hg (P<0.03)⁽¹⁷³⁾. Soon after, Webb et al. investigated this topic further using beetroot juice 553 (containing approximately 1400 mg inorganic nitrate)⁽³³⁾. Results from Webb et al. showed a peak 554 reduction in systolic blood pressure of $10.4 \pm 3 \text{ mm Hg}$ (P<0.01), a reduction in diastolic blood 555 pressure of 8.1 \pm 2.1 mm Hg (P<0.01) and mean arterial pressure reduction of 8.0 \pm 2.1 mm Hg 556 (P<0.01), thus indicating that significant blood pressure reductions are possible with the acute 557 consumption of dietary inorganic nitrate in healthy subjects ⁽³³⁾. A notion which has been further 558 559 supported by a recently conducted systematic review and meta-analysis which found inorganic nitrate and beetroot juice consumption were associated with greater changes in systolic blood 560 pressure (-4.4 mm Hg (95% CI: -5.9, -2.8); P<0.001) than diastolic blood pressure (-1.1 mm Hg 561 (95% CI: -2.2, 0.1); P=0.06)⁽¹⁷⁴⁾. However it is important to note that these findings have not been 562 consistent across the literature, as a few recently conducted randomised controlled trials have found 563 inorganic nitrate consumption from either beetroot juice or from a high nitrate diet (rich in green 564 leafy vegetables) for 1-2 weeks had little/no effect on the blood pressure of study subjects (57; 175; 565 ¹⁷⁶⁾. The exact cause of this variation across studies remains unclear, yet could be due to 566 methodological differences including the study population (e.g. healthy subjects vs. hypertensive 567 subjects) or the conditions in which NO_x was consumed (e.g. food vs supplement, dosing or altered 568 environmental conditions such as exercise stress). Nevertheless, this question remains unclear and 569 will require further investigation, in order to better understand the usefulness of dietary/inorganic 570 571 nitrate/nitrite within the general population.

572

While the acute effects of dietary inorganic nitrate on blood pressure has been extensively 573 574 investigated, very few studies have investigated long-term effects. Sobko et al. investigated the effects of a traditional Japanese diet on blood pressure which provided approximately 1140 mg of 575 nitrate per day for a 10 day period ⁽²³⁾. The Traditional Japanese diet lead to a lower diastolic blood 576 pressure than seen in the non-Japanese diet group (71.3 \pm 7.9 vs 75.8 \pm 7.8, P=0.0066), indicating 577 578 that dietary inorganic nitrate consumption for longer-periods of time may have some blood pressure 579 lowering effects in healthy people, however a 10 day intervention can hardly be classified as a long-580 term intervention ⁽²³⁾. In another four week intervention Kapil et al. assigned hypertensive patients 581 to receive a daily dose of either 250 mL of beetroot juice or placebo (nitrate depleted beetroot juice)⁽²⁹⁾. Notably, Kapil et al. found daily dietary nitrate supplementation to significantly reduced 582 mean clinic blood pressure (7.7/2.4 mm Hg (3.6-11.8/0.0-4.9), P<0.001, P=0.05), mean 24-hour 583

ambulatory blood pressure (7.7/5.2 mm Hg (4.1-11.2/2.7-7.7), P<0.001 for both) and mean home
blood pressure (8.1/3.8 mm Hg (3.8-12.4/0.7-6.9), P<0.001, P<0.01) ⁽²⁹⁾.

Currently, the longest intervention study conducted in this area is a 10 week intervention trial from 586 DeVan et al⁽¹²⁵⁾. In this study, healthy 50-79 year old subjects were recruited to consume either 0 587 mg, 80 mg or 160 mg of sodium nitrite per day for a 10 week period ⁽¹²⁵⁾. Results indicated no 588 significant changes in blood pressure at week 10 compared with baseline blood pressure values, 589 however a significant time by treatment effect for carotid diameter in the nitrite groups was 590 detected, as well as improved endothelial function of the brachial artery, suggesting improved 591 vascular function with chronic inorganic nitrite supplementation despite a lack of an effect seen 592 with blood pressure⁽¹²⁵⁾. However, it is worth noting that the only perspective cohort study on this 593 topic conducted by Golzarand et al. found that a higher dietary intakes of nitrate containing 594 vegetables (~427.6 g/day) in normotensive individuals may have a protective effect against the 595 development of hypertension (Highest tertile of nitrate containing vegetables, OR: 0.63 (0.41-0.98), 596 P=0.05)⁽¹⁷⁷⁾. 597

Endothelial dysfunction is one of the key early events involved in the development of 598 atherosclerosis ⁽¹⁷⁸⁾. Flow mediated dilatation is commonly used as a measure of endothelial 599 function as reduced flow mediated dilatation is an indicator of endothelial dysfunction (caused by 600 601 reduced NO bioavailability) and has been associated with increased severity and duration of blood pressure elevations (179). More recently, dietary inorganic nitrate interventions have been shown to 602 significantly improve flow mediated dilatation in healthy and hypertensive humans consuming 603 spinach, beetroot juice or sodium nitrate capsules ^(29; 168; 180; 181). Joris et al. tested the effects of 604 beetroot juice (containing approximately 500 mg nitrate) with a dietary load of fat (56.6 g fat) in 605 overweight and obese subjects (BMI: $30.1 \pm 1.9 \text{ kg/m}^2$)⁽¹⁸²⁾. While the control drink group saw 606 impaired flow mediated dilatation with dietary fat intake, the consumption of beetroot juice 607 appeared to attenuate this impairment (Beetroot juice: $-0.37 \pm 2.92\%$ vs Control: $-1.56 \pm 2.9\%$, 608 P=0.03)⁽¹⁸²⁾. Additionally, flow mediated dilatation has been shown to be reduced by 609 approximately 40% after vascular ischemia, however Ingram et al. has demonstrated that sodium 610 nitrite pre-conditioning (providing nitrite dose prior to ischemic event) will prevent ischemic 611 reperfusion injury by preventing reductions in flow mediated dilatation and endothelial dysfunction 612 ⁽¹⁸³⁾. Similar findings have been reported by Kapil et al. and Webb et al. with beetroot juice pre-613 conditioning, indicating that higher plasma NO_x concentrations achieved by inorganic NO_x 614 consumption may have a role for improving cardiovascular outcomes post vascular ischemic events 615 (29; 33) 616

- 617 In addition to flow mediated dilatation, CACs have been identified as an important indicator of
- 618 vascular endothelial function, as they have a critical role in vascular repair ⁽¹⁸⁴⁾. The number of
- CACs have also been shown to predict the occurrence of cardiovascular disease and death ⁽¹⁶⁸⁾.
- 620 Therefore it is of interest to note that Heiss et al. have indicated an important role for dietary nitrate
- 621 for increasing CACs, showing that a single dose of sodium nitrate (12.7 mg/kg body weight) can
- 622 double the number of CACs 1-2 hours post nitrate ingestion $^{(168)}$.
- Pulse wave velocity and augmentation index are accepted measurements of arterial stiffness and 623 atherosclerosis, to which higher readings are associated with increased cardiovascular disease risk 624 ^(185; 186). The role for dietary inorganic nitrate in preventing arterial stiffness has been established, as 625 Kapil et al. found a 4 week beetroot juice intervention to reduce pulse wave velocity and 626 augmentation index in hypertensive subjects ⁽²⁹⁾. Zamani et al. also saw a significantly reduced 627 augmentation index with beetroot juice consumption in patients with symptomatic heart failure 628 (Beetroot juice: $132.2 \pm 16.7\%$; Placebo: $141.2 \pm 21.9\%$; mean change $-9.1 \pm 15.4\%$; P=0.03) ⁽¹⁸⁷⁾. 629 Rammos et al. investigated the effect of a 4 week sodium nitrate supplementation trial in elderly 630 volunteers with mild hypertension, and found that vascular stiffness was significantly improved in 631 the nitrate supplemented volunteers ⁽¹⁸⁸⁾. This is a very significant finding given that vascular 632
- 633 stiffness tends to naturally increase with age (189).
- In an RCT conducted by Jones et al. participants prone to MI and undergoing primary percutaneous coronary intervention (non-surgical intervention to treat stenosis) were administered with either a high-dose bolus injection of NaNO₂⁻ (1.8 μ mol) or NaCl placebo⁽¹⁹⁰⁾. The nitrite group experienced a significantly (P=0.05) improved myocardial savage index (established indicator of cardio protective benefit) relative to placebo⁽¹⁹⁰⁾. In addition, a sub-set of participants which exhibited a blocked blood vessel experienced a 19% reduction in infarct size with nitrite treatment compared to placebo⁽¹⁹⁰⁾. A one-year follow-up of study participants also found the nitrite group experienced a
- 641 significant reduction in major adverse cardiac events (NaNO₂⁻: 2.6% vs NaCl: 15.8%, P=0.04)⁽¹⁹⁰⁾.

642 Conclusion

643 Cardiovascular disease remains the major killer from any disease across the developed world. Currently the available evidence indicates a role for dietary nitrate for improving cardiovascular 644 645 disease risk factors, a highly valuable finding given that dietary nitrate from beetroot and green leafy vegetables could represent a relatively simple and cost effective treatment/preventative 646 647 strategy for reducing CVD and its sequelae. However, at present it remains unclear whether incidence of cardiovascular disease morbidity or mortality can be reduced with long-term dietary 648 649 intakes of inorganic nitrate, as such evidence investigating this question directly has not yet been published. At present, there is an overwhelming need for epidemiological research to be conducted 650

651	to identify the potential long-term effects of sustained inorganic nitrate and nitrite consumption on
652	the development of cardiovascular disease and its consequences.
653	
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1177 Tables

1178 Table 1. Permissions for nitrate and nitrite in Food Products

Product	Additive	Maximum Permitted Level (mg/kg
	Nitrite Salt	150 (90)
	Nitrate Salt	50 (85; 90)
Commercially Sterile Canned Dried Meat	Nitrite Salt	50 - 150 (85; 91)
	Nitrite Salt	125 (90)
	Nitrate Salt	150 (85)
	Nitrite Salt	125 - 200 ^(85; 90)
	Nitrate Salt	175- 500 ^(85; 92) 125 - 175 ^(85; 90)
	Nitrite Salt Nitrate Salt	125 - 175 (90; 91) 150 - 300 (90; 91)
	Nitrite Salt	150 - 500 (90)
	Nitrate Salt	500 (85; 90)
Nitrate salt: Potassium Nitrate and Sodium Nitrate. Nitrite salt: Po		
	Jussium Plurice und Sourd	in the let

1203	Table 2. Vegetable sources of nitrate and nitrite with estimated nitrate and/or nitrite contents.
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Vegetable Type	Nitrate Content (mg/kg) Mean (range)	Nitrite Content (mg/kg) Mean (range)
Rocket	3624 (1550-7316) ^(111; 191)	NA
Turnip Greens	3467 ^(192; 193; 194)	NA
Spinach	2485 (2-6700) ^{(22; 79; 85; 111; 191; 193; 194; 195; 196; 197; 198; 199;}	15 (ND-162) ^(22; 85; 200; 202; 203; 205)
	200; 201; 202; 203; 204; 205)	
Swiss chard	2363 ⁽¹⁹⁹⁾	NA
Turnip	2174 (10-4800) ^(111; 194; 195; 197; 201)	NA
Rhubarb	$1999\ (55\text{-}6500)^{(191;\ 193;\ 194;\ 196;\ 197;\ 201;\ 204)}$	NA
Celery	1964 (19-5300) ^(85; 191; 193; 194; 195; 196; 197; 198; 199; 201; 203)	2.5 (ND-6) ^(85; 191)
Beetroot	1992 (100-8100) ^(85; 111; 193; 194; 195; 196; 197; 198; 199; 201; 203; 204; 206; 207; 208)	1.7 (ND-110) ^(85; 199; 203; 209)
Chinese Cabbage	1855 (111-8050) ^(201; 202; 206; 208; 210)	0.9 (ND-14.3) ^(206; 208)
Radish	1773 (60-9000) ^(111; 191; 193; 194; 195; 196; 201)	NA
Lettuce	$1689\ (10\text{-}13000)^{(79;\ 85;\ 111;\ 191;\ 193;\ 194;\ 195;\ 196;\ 197;\ 198;\ 199;}$	0.8 (ND-5) ^(85; 203; 205; 206; 208)
	201; 202; 203; 204; 205; 206; 208; 209)	· (202)
Watercress	$1640 (890-2790)^{(203)}$	2.5 (ND-5) ⁽²⁰³⁾
Buk Choy	1620 (1023-3098) ⁽²⁰²⁾	$20 (0.09-30)^{(202)}$
Kale/ Mustard Greens	1318 (19-5500) ^(22; 191; 192; 193; 194; 197; 205)	$(0.03-0.64)^{(22;205)}$
Silver beet	1255 (190-1770) ^(203; 209)	2.5 (ND-5) ^(203; 209)
Endive	975 (10-3800) ^(194; 199)	NA
Broccoli	$793 \ (ND\text{-}2300)^{(22; 85; 193; 194; 196; 197; 198; 199; 203; 204)}$	3 (ND-110) ^(22; 85; 203)
Cabbage	756 (1-3100) ^(85; 193; 194; 195; 196; 197; 198; 199; 201; 203; 204; 207; 208; 209; 210)	0.8 (ND-26) ^(85; 203; 208)
Cauliflower	547 (ND-4500) ^(191; 193; 194; 195; 196; 197; 198; 199; 201)	NA
Mixed Salad	540 (80-821) ^(22; 111; 191; 201)	1.3(22)
Eggplant	479 (31-1500) ^(191; 194; 195; 198; 199)	NA
Leek	399 (56-841) ^(111; 195)	NA
Pumpkin / Squash	389 (ND-2200) ^(85; 191; 194; 195; 196; 197; 198; 199; 201; 203)	6 (ND-194) ^(85; 203)
Green Onion	366 (4-1676) ^(111; 201)	NA
Fennel	363 ⁽¹⁹⁹⁾	NA
Green Beans	315 (6-1100) ^(85; 111; 193; 195; 197; 199; 208)	7 (0.16-57) ^(85; 208)
Cucumber	$184\ (1\text{-}1236)^{(85;\ 111;\ 191;\ 194;\ 195;\ 198;\ 199;\ 208;\ 209;\ 210)}$	3 (ND-1164) ^(85; 208)
White Potato	184 (ND-5521) ^(22; 85; 111; 191; 193; 194; 195; 196; 197; 198; 201; 203; 207; 208; 209; 210)	1 (ND-10.3) ^(22; 85; 203; 208)
Carrot	182 (ND-2800) ^{(22; 85; 111; 191; 193; 194; 195; 196; 197; 198; 199;} 201; 203; 204; 205; 207; 208)	0.7 (ND-7.5) ^(22; 85; 203; 205; 208)
Garlic	163 (1-462) ^(111; 191; 199)	NA
Lima Beans	160 (54-310) ^(193; 195; 198)	NA
Brussels Sprouts	118 (ND-170) ⁽¹⁹⁴⁾	NA
Onion	100 (ND-2300) ^(85; 191; 194; 195; 196; 199; 201)	0.5 (ND-2.2) ⁽⁸⁵⁾
Mushroom	92 (ND-400) ^(85; 191; 194)	NA
Asparagus	84 (13-700) ^(194; 196; 198)	NA
Tomato	71 (ND-392) ^{(22; 85; 111; 191; 193; 194; 195; 196; 198; 199; 201; 204;}	0.6 (ND-13) ^(22; 85; 208)
Toniato	207; 208; 209; 210)	0.0 (10 10)
Sweet Potato	55 (ND-66) ^(191; 193; 194; 195; 198)	NA
Peas	32 (ND-124) ^(85; 191; 193; 194; 195; 198; 199)	(ND-22) ⁽⁸⁵⁾
	30 (9-68) ^(195; 198)	(ND-22) ^(a) /NA
Dry Beans	30 (ND-45) ^(85; 195; 198)	
Corn	30 (ND-45) ^(64, 154, 165) 30 ⁽¹⁹⁹⁾	(ND-7.5) ⁽⁸⁵⁾
Artichoke	30 ⁽¹³³⁾ 22 (21-23) ⁽⁸⁵⁾	NA NA
Preserved Olives		NIA

Vegetable Type	Nitrate Content (mg/kg) Mean (range)	Nitrite Content (mg/kg) Mean (range)		
Baked Beans	17 (ND-23) ⁽⁸⁵⁾	1.7 (ND-7.5) ⁽⁸⁵⁾		
Data is combined nitrate and nitrite a available.	estimates from various published papers, government	documents and reviews. ND: Not Detected. NA: I		
Table 3. Meat based sources of nitrate and nitrite with estimated nitrate and/or nitrite contents.				
Meat Type				
Meat Type	Nitrate Content (mg/kg)	Nitrite Content (mg/kg)		
Meat Type	Nitrate Content (mg/kg) Mean (range)	Nitrite Content (mg/kg) Mean (range)		
Meat Type Salami				
	Mean (range)	Mean (range)		
Salami	Mean (range) 94 (ND-450) ^(85; 202; 203; 211; 212; 213)	Mean (range) 31 (ND-108) ^(85; 202; 203; 211; 212; 213)		
Salami Bolonga	Mean (range) 94 (ND-450)(85: 202; 203; 211; 212; 213) 65 (4-98)(211; 214; 215) 64 (8-81)(22; 85; 202; 203)	Mean (range) 31 (ND-108) ^(85; 202; 203; 211; 212; 213) 14 (ND-55) ^(211; 214; 215; 216)		
Salami Bolonga Frankfurt/Hot dog	Mean (range) 94 (ND-450)(^{85; 202; 203; 211; 212; 213)} 65 (4-98)(^{211; 214; 215)} 64 (8-81)(^{22; 85; 202; 203)}	Mean (range) 31 (ND-108) ^(85; 202; 203; 211; 212; 213) 14 (ND-55) ^(211; 214; 215; 216) 39 (0.5-95) ^(22; 85; 202; 203)		
Salami Bolonga Frankfurt/Hot dog Shelf-Stable, Canned Cured N	$\begin{array}{c} \mbox{Mean (range)} \\ \mbox{94 (ND-450)(85; 202; 203; 211; 212; 213)} \\ \mbox{65 (4-98)^{(211; 214; 215)}} \\ \mbox{64 (8-81)^{(22; 85; 202; 203)}} \\ \mbox{Meat} \mbox{63 (ND-840)^{(211; 212; 214)}} \end{array}$	Mean (range) 31 (ND-108) ^(85; 202; 203; 211; 212; 213) 14 (ND-55) ^(211; 214; 215; 216) 39 (0.5-95) ^(22; 85; 202; 203) 31 (ND-19) ^(211; 212; 214) 33 (ND-940) ^(91; 202; 211; 214; 216; 217; 218; 219; 220)		
Salami Bolonga Frankfurt/Hot dog Shelf-Stable, Canned Cured M Sausages	$\begin{array}{c} \mbox{Mean (range)} \\ 94 (ND-450)^{(85; 202; 203; 211; 212; 213)} \\ 65 (4-98)^{(211; 214; 215)} \\ 64 (8-81)^{(22; 85; 202; 203)} \\ \mbox{Meat} & 63 (ND-840)^{(211; 212; 214)} \\ 58 (15-240)^{(85; 202; 211; 214; 217; 218)} \end{array}$	Mean (range) 31 (ND-108) ^(85; 202; 203; 211; 212; 213) 14 (ND-55) ^(211; 214; 215; 216) 39 (0.5-95) ^(22; 85; 202; 203) 31 (ND-19) ^(211; 212; 214) 33 (ND-940) ^(91; 202; 211; 214; 216; 217; 218; 219; 220)		

Bacon	42 (ND-310)(-1, 44, 54, 54, 54, 54, 54, 54, 54)	29 (ND-430) ^{(,,,,,,,,}
		219; 222; 223; 224)
"Luncheon Meat"	32 (<10-70) ^(85; 203; 215)	31 (ND-130) ^(85; 203; 215)
Pork	21 (ND-19) ^(22; 215)	(ND-8) ^(22; 215)
Corned Beef	14 (4-36) ^(203; 215)	3 (ND-8) ^(203; 215)
Minced Beef	12 (ND-24) ^(202; 203)	NA

1212 Data is combined nitrate and nitrite estimates from various published papers, government documents and reviews. ND: Not Detected. NA: Data not available.

1213

1214

1215 Table 4. Fruit sources of nitrate and nitrite with estimated nitrate and/or nitrite contents.

Fruit Type	Nitrate Content (mg/kg) Mean (range)	Nitrite Content (mg/kg) Mean (range)
Melon	325 (38-600) ^(194; 195; 196; 199; 201)	NA
Strawberries	172 (96-233) ⁽⁸⁵⁾	18 (8-80) ⁽⁸⁵⁾
Banana	76 (45-200) ^(22; 85)	2 (ND-11) ^(22; 85)
Apple	20 (ND-56) ⁽⁸⁵⁾	(ND-7.5) ⁽⁸⁵⁾
Grapes	19 (ND-52) ⁽⁸⁵⁾	10 (ND-19.4) ⁽⁸⁵⁾
Sultanas	16 (9-22) ⁽⁸⁵⁾	0.8 (ND-5.5) ⁽⁸⁵⁾
Peach	10 (7-18) ⁽⁸⁵⁾	17 (ND-22) ⁽⁸⁵⁾
Orange	9 (ND-21) ^(22; 85)	0.2 (ND-7.5) ⁽⁸⁵⁾
Mango	9 (ND-12) ⁽⁸⁵⁾	6 (ND-15) ⁽⁸⁵⁾
Watermelon	8 (7-18) ⁽⁸⁵⁾	(ND-16.4) ⁽⁸⁵⁾
Pineapple	7 (ND-12) ⁽⁸⁵⁾	17 (10-22) ⁽⁸⁵⁾

1216 Data is combined nitrate and nitrite estimates from various published papers, government documents and reviews. ND: Not Detected. NA: Data not 1217 available.

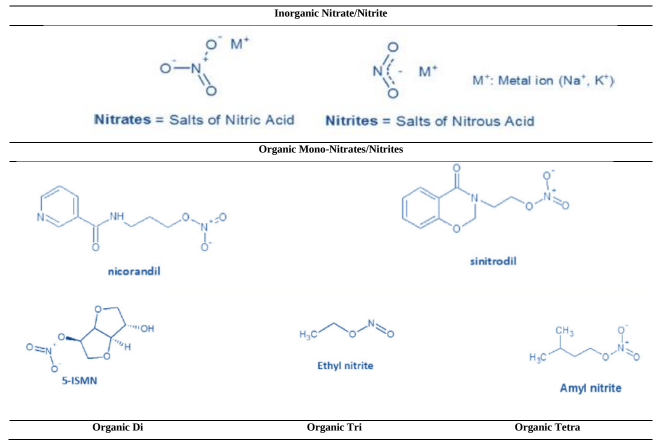
1219 Table 5. Nitrate and nitrite containing herbs with estimated nitrate and/or nitrite contents.

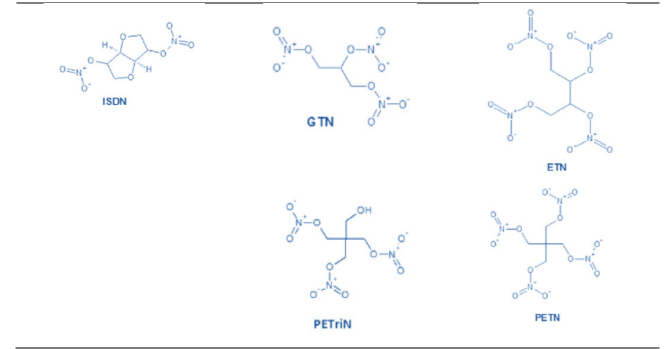
Herb Type	Nitrite Content (mg/kg) Mean (range)	Nitrite Content (mg/kg)	
		Mean (range)	
Dill	2590 (2236-3267) ^(200; 201)	102(200)	
Parsley	1304 (ND-4467) ^(85; 194; 195; 196; 200; 201)	(ND-94) ^(85; 200)	
Tea	3 (2-3) ⁽⁸⁵⁾	(ND-0.3) ⁽⁸⁵⁾	

¹²¹⁸

- 1220 Data is combined nitrate and nitrite estimates from various published papers, government documents and reviews. ND: Not Detected. NA: Data not
- 1221 available.
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1227 Table 6. Chemical structure of inorganic nitrate/nitrite compared with organic mono-, di-, tri- and tetra nitrates/nitrites.





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1228Table from Omar, Artime and Webb, 2012 (83).
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