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Claire R. Palmer
Edith Cowan University

Lauren C. Blekkenhorst
Edith Cowan University

Joshua R. Lewis
Edith Cowan University

Natalie C. Ward

Carl J. Schultz

See next page for additional authors

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Authors

Claire R. Palmer, Lauren C. Blekkenhorst, Joshua R. Lewis, Natalie C. Ward, Carl J. Schultz, Jonathan M. Hodgson, Kevin D. Croft, and Marc Sim

1 **Review article**

2 Quantifying dietary vitamin K and its link to cardiovascular health: a narrative review.

3 Claire R Palmer^{1,2}, Lauren C Blekkenhorst^{2,3}, Joshua R Lewis^{2,3,4}, Natalie C Ward^{3,5}, Carl J
4 Schultz^{3,6}, Jonathan M Hodgson^{2,3}, Kevin D Croft¹, Marc Sim^{2,3}

5 ¹ School of Biomedical Sciences, The University of Western Australia, Perth, Western
6 Australia, Australia

7 ² School of Health and Medical Sciences, Edith Cowan University, Perth, Western Australia,
8 Australia

9 ³ School of Medicine, Faculty of Health and Medical Sciences, The University of Western
10 Australia, Perth, Western Australia, Australia

11 ⁴ Centre for Kidney Research, Children's Hospital at Westmead, School of Public Health,
12 Sydney Medical School, The University of Sydney, Sydney, New South Wales, Australia

13 ⁵ School of Public Health & Curtin Health Innovation Research Institute, Curtin University,
14 Perth, Western Australia, Australia

15 ⁶ Department of Cardiology, Royal Perth Hospital, Perth, Western Australia, Australia

16 **Corresponding author**

17 Marc Sim

18 School of Medical and Health Sciences, Edith Cowan University

19 Joondalup, 6027, Western Australia, Australia

20 Email: marc.sim@ecu.edu.au

21

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23 Abstract

24 Cardiovascular disease is the leading cause of death and disability worldwide. Recent work
25 suggests a link between vitamin K insufficiency and deficiency with vascular calcification, a
26 marker of advanced atherosclerosis. Vitamin K refers to a group of fat-soluble vitamins
27 important for blood coagulation, reducing inflammation, regulating blood calcium metabolism,
28 as well as bone metabolism, all of which may play a role in promoting cardiovascular health.
29 Presently, there is a lack of a comprehensive vitamin K database on individual foods, which
30 are required to accurately calculate vitamin K1 and K2 intake for examination in
31 epidemiological studies. This has likely contributed to ambiguity regarding the recommended
32 daily intake of vitamin K, including whether vitamin K1 and K2 may have separate, partly
33 overlapping functions. This review will discuss the presence of: (i) vitamin K1 and K2 in the
34 diet; (ii) the methods of quantitating vitamin K compounds in foods; and (iii) provide an
35 overview of the evidence for the cardiovascular health benefits of vitamin K in observational
36 and clinical trials.

37 Key words

38 Vitamin K, Database, Phylloquinone, Menaquinone, Vascular calcification, Cardiovascular
39 health

40

41 **Introduction**

42 Vitamin K refers to a group of fat-soluble vitamins important for blood coagulation and has
43 been linked to other biological processes including blood calcium metabolism, as well as bone
44 and vascular health.¹ The two main vitamin K forms are vitamin K1 (phylloquinone; PK) and
45 vitamin K2 (menaquinones; MK). The few studies that have explored the link between vitamin
46 K1 and/or K2 with cardiovascular outcomes have reported conflicting results.¹ Despite the
47 potential importance of vitamin K1 and/or K2 for human health, the adequate intake (AI)
48 recommendations proposed by numerous public health organisations typically only consider
49 vitamin K1. This is likely a result of limited information relating to vitamin K2.² Specifically,
50 there is currently a lack of international comprehensive databases on the vitamin K content of
51 individual foods, which are required to accurately calculate vitamin K1 and/or K2 intake.² This
52 review discusses the: (i) presence of vitamin K1 and K2 in the diet; (ii) the methods of
53 quantitating vitamin K compounds in foods; and (iii) provides an overview of the evidence for
54 the cardiovascular health benefits of vitamin K in observational and clinical trials. We also
55 propose that the development of a comprehensive national vitamin K databases of individual
56 food items is critical to advance our understanding of vitamin K nutrition for a range of health
57 outcomes including cardiovascular health.

58 **Vitamin K: phylloquinone and menaquinones**

59 Vitamin K is a group of structurally and functionally analogous fat-soluble vitamins, which act
60 as cofactors in the post-translational γ -carboxylation of vitamin K-dependant proteins (VKDP).
61 Vitamin K compounds are characterised by a 2-methyl-1,4-naphthoquinone backbone also
62 called menadione and a lipophilic isoprenoid side chain attached at carbon three (*Figure 1*).
63 Two types of vitamin K exist naturally: vitamin K1 (PK) and vitamin K2 (MK).³ Phylloquinone
64 has a side chain analogous to the phytyl side chain of chlorophyll; four isoprenoid residues of
65 which three are saturated. It is synthesised by all photosynthetic plants and algae, as it is a

66 functional component in photosynthesis.⁴ Vitamin K2 (MK) is composed of a group of
67 isoprenologs where the side chain is a polymer of isoprenoid units ranging between four to
68 thirteen repeats. MKs are named according to the number of repeating units,⁴ with derivatives
69 including MK-4 to MK-13. For example, MK-4 has four unsaturated isoprenoid units (*Figure*
70 *1*). Despite having analogous structures the origins of MKs differ. MK-4 is synthesised from
71 PK and menadione by animals and humans.² All other MKs are synthesised by anaerobic
72 bacteria, which includes species present in the human microbiome.⁴ Bacterial MKs are
73 important in metabolic respiration acting as components of the electron transport chain.⁵ The
74 synthetic compound, menadione, is also referred to as vitamin K3 and is often added to animal
75 feed.⁶

76 *Dietary vitamin K*

77 Phylloquinone is present in the majority of foods including vegetables, fruits, nuts, bread,
78 cereals, oil, meat and animal products; however, quantities vary greatly.^{7, 8} The highest
79 concentrations of PK found in green leafy vegetables, due to its role in photosynthesis.
80 Concentrations in these vegetables range between 400-700 $\mu\text{g}/100\text{ g}$.⁴ Vegetable oils, such as
81 olive, canola and soybean, also contain considerable amounts of PK. These oils are widely used
82 in margarine, salad dressings, and the preparation of foods, such as bread. The contribution of
83 oils as part of commercially consumed foods towards total dietary vitamin K intake may be
84 greater than initially thought. For example, in the US, commercially prepared foods and mixed
85 dishes were the second highest contributors to PK intake, after vegetables.⁹ The widespread
86 use of plant products in food production and their consumption by animals explains the
87 presence of PK in disparate food groups. The PK content in the Western diet makes up the
88 majority of vitamin K intake. It constitutes up to 90% of dietary vitamin K intake, of which 60-
89 80% is derived from vegetables.^{4, 10, 11}

90 Menaquinones are found in meat, dairy and fermented foods.² MK-4 is synthesised by the
91 conversion of consumed PK or menadione in human or animal tissues. Therefore, dietary MK-
92 4 is typically found in animal tissue and products, such as eggs or dairy.² The tissue or organ
93 of an animal, the type of animal and country of origin (possibly related to feed) all affects the
94 MK-4 content in food products.² For example, goose liver is a major storage tissue of vitamin
95 K compounds and has the highest MK-4 content of any tissue (370 µg/100 g).³ In comparison,
96 goose leg has only 60 µg/100 g of MK-4.³ Alternatively, chicken breast has more MK-4 than
97 duck breast; 8.9 vs 3.6 µg/100 g, respectively.³ Poultry and pork, have especially high MK-4
98 content due to menadione in their feed, which is converted into MK-4.^{2, 12} The MK-4 content
99 of various poultry products vary greatly.² Presently, the most comprehensive tool to calculate
100 the MK-4 content of food is provided by the United States Department of Agriculture (USDA)
101 Food Composition Databases.¹³ However, there is limited information on the MK-4 content in
102 some commonly consumed meats, such as lamb, veal, and goat.¹⁴ Other MKs (MK5-12) are
103 synthesised by anaerobic bacteria⁴; however, information on their approximate content in food
104 products are scarce.

105 Foods fermented with the MK-synthesising bacteria are abundant in long-chain MKs. For
106 example, natto (fermented soybeans) has been reported to have the highest MK-7 content of
107 any food, containing 936 µg/100 g.¹⁵ The type of MKs produced is dependent on the type of
108 bacteria and temperature conditions.⁵ Notably, bacteria that produce MK-9 also produce MK-
109 8, but in four-fold lower amounts. Thus, MK-9 and MK-8 co-occur in foods that are fermented.
110 Other MKs are produced independent of each other.¹⁶ Not all fermented foods contain MKs.
111 For example, yeast has lost the enzymes required for MK synthesis,^{4, 5} thus bread does not
112 contain MKs.² Fermented dairy products, such as cheese, milk and yogurt, are significant
113 sources of long-chain MKs.^{2, 17} The MK content varies by the percentage of fat present in dairy.

114 For example, the MK content of full fat cheddar cheese is five times higher than the total MK
115 content of reduced-fat cheddar (281 $\mu\text{g}/100\text{ g}$ vs. 49 $\mu\text{g}/100\text{ g}$, respectively).¹⁷

116 Few studies have analysed the MK content in the Western diet. From this limited work, MKs
117 have been reported to constitute approximately 10% of dietary vitamin K intake.^{4, 10, 11} Of note,
118 the major MK form is MK-4, which contributes 25-40% of the total MK in the diet.^{10, 11} After
119 MK-4, the most nutritionally abundant MKs are MK-9, MK-8 and MK-7.^{10, 18} However, as the
120 MK content in individual foods has not been systematically determined to the degree of PK,²
121 this may lead to errors in these estimates. Studies have highlighted meat and dairy products as
122 the best sources of MKs.³ For example, a German study reported that 60% of all MKs came
123 from dairy products, with cheese consumption contributing 43% of total MK intake.¹⁰
124 Considering, the European diet is typically rich in cheese, it is thought to be a major contributor
125 of MK intake in such populations.² Presently, the most comprehensive resources for the
126 different MK content from a variety of animal and plant-based foods is from the Netherlands.³
127 Besides the aforementioned Dutch work, there is relatively little data for the MK content in the
128 diets of other countries. This is likely due to the absence of comprehensive databases of MK
129 content in individual foods, which are required to quantify MK intake at individual and
130 population levels.²

131 *Bioavailability of vitamin K*

132 Bioavailability is the rate and extent by which a nutrient is absorbed and becomes available to
133 the target tissue.¹⁹ Alternatively, bioaccessibility relates to the amount of compound that is
134 released from its matrix in the gastrointestinal tract and is available for absorption.¹⁹ Precise
135 quantitative assessments of vitamin K bioavailability in humans is challenging due to
136 unquantified tissue conversion of PK to MK-4 and the contributions of the gut microbiota.²⁰
137 Both PK and MKs are absorbed from the intestine using the same mechanisms as other fat
138 soluble vitamins and lipids.⁴ Apart from this shared absorption mechanism there are differences

139 in the bioavailability and bioaccessibility of vitamin K compounds.³ For example, previous
140 work examining the absorption rate of PK and MK in a food matrix reported that MK
141 absorption from natto (Japanese fermented soybean) is approximately 10 times greater
142 compared to PK absorption from spinach. Of note, peak serum values for both PK and MK
143 appear to occur at the ~6 h post-prandial mark. It was also reported that bioavailability of PK
144 from foods is significantly reduced by its poor bioaccessibility from the food matrix.³
145 Specifically, only 5-10% of PK is absorbed from cooked vegetables; however, this can be
146 increased slightly by the presence of dietary lipids.³ In contrast, PK absorption from oils and
147 supplements have much higher absorption rates (~200-400%) compared to vegetables.^{3, 21, 22}
148 This indicates that PK as a nutrient can be well absorbed, but the vegetable food matrix may
149 present a barrier to absorption. Compared to PK, the limited amount of studies conducted on
150 long-chain MKs suggest they may have higher bioaccessibility from dietary sources³.
151 Specifically, the absorption of long-chain MKs (MK-7, MK-8 and MK-9) from natto, cheese
152 and egg yolk is close to 100%.³ Furthermore, these long-chain MKs have a longer half-life. For
153 example, in a study where 6 male volunteers were given 2 μmol of PK, MK-4 and MK-9, MK-
154 9 was still detectable in the serum at 48 h, while PK and MK-4 returned to baseline after 24
155 h.²³ Notably, the long half-life of MK-9 may not necessarily indicate increased bioavailability,
156 but instead non-preferential utilisation by tissues compared to PK and MK-4. A long half-life
157 may also indicate that long-chain MKs may be of particular importance for extrahepatic
158 tissues.²⁴ Despite the aforementioned work, the bioavailability of vitamin K compounds from
159 different food matrixes remain largely unknown. Nevertheless, current evidence suggest that
160 long-chain MKs may be more important than previously thought for vitamin K status, due to
161 the almost complete absorption of long-chain MKs and poor absorption of PKs from its main
162 dietary sources.²⁵

163

164 *Non-dietary sources of menaquinones*

165 Despite, the large proportion of animal-based foods known to contain MKs, not all of the MKs
166 come from dietary sources. There are two known alternative sources, including PK conversion
167 into MK-4, and MK synthesis by the gut microbiota.^{4, 26} The full extent of the conversion of
168 PK to MK-4 is unknown, but in the last decade progress has been made in understanding the
169 mechanism. Specifically, intestinal PK is cleaved in enterocytes to produce menadione (often
170 referred to as vitamin K3), which is prenylated by UbiA prenyltransferase-containing domain
171 1 (UBIAD1) to produce MK-4.^{4, 26} A recent study in rats demonstrated that only 0.1% of oral
172 PK was converted into MK-4, and 2% converted into menadione.²⁷ The extent of this
173 conversion in humans remains unclear, but it is estimated to be 5-25% of PK intake.²⁸
174 Therefore, MK-4 biosynthesis from PK may be a large source of MKs, but detailed human
175 studies are needed to examine this further. In addition, the contribution of microbial-
176 synthesised MKs to human nutrition is not fully understood. The majority of MKs in the gut
177 are synthesised by *Bacteroides*-one of the largest intestinal genera.^{29, 30} *Bacteroides* primarily
178 produce MK-10 and MK-11, with small amounts of MK-7 to MK-9.^{29, 31} There is a large pool
179 of MKs in the gut, with previous work suggesting an average of 1.8 mg of vitamin K2 in the
180 intestinal tract of humans.³² Despite such reserves of vitamin K2 being over 20 times the AI
181 for adult males, the bioavailability of these MKs is thought to be limited.³³ Specifically, vitamin
182 K compounds are absorbed in the small intestine in a process requiring bile salts and pancreatic
183 enzymes, which are absent where the majority of bacteria colonise the colon.³⁴ Additionally,
184 the majority of microbial-synthesized MKs are bound by bacterial membranes and thus are not
185 available for absorption.² Therefore, while a large amount of MKs are produced by the gut
186 microbiota, their poor bioavailability at the point of synthesis limits their contribution to human
187 vitamin K nutrition.^{31, 33} Presently, the exact bioavailability of microbial-synthesised MKs is
188 unknown, and more bioavailability studies are required to discern their role in human nutrition.

189 *Dietary recommendations for vitamin K*

190 Current dietary vitamin K recommendations are typically based on the intake of PK sufficient
191 to maintain coagulation.²⁰ This is problematic as the tissue-requirements of extrahepatic
192 vitamin K-dependant proteins for PK differs from the requirement of hepatic vitamin K-
193 dependant proteins.² Therefore, dietary intakes which are sufficient to maintain coagulation
194 may be inadequate in optimising other aspects of health that might be impacted by vitamin K
195 nutrition.^{2, 28} However, any attempts to update the AI values for vitamin K1 or include vitamin
196 K2 would be hampered by a lack of dose-response studies and evidence on the occurrence,
197 absorption, function and content of MKs in the body or organs.^{28, 35, 36} Difficulties in
198 quantitating the contribution of MKs synthesised by the gut microflora and tissue conversion
199 of PK are additional challenges when defining dietary requirement.²⁰ Furthermore, a recent
200 report by the European Commission highlighted the uncertainties in country-wide estimates of
201 vitamin intake data, which are caused by ‘borrowing’ vitamin K values from other countries
202 and substituting vitamin K values for individual food items for similar foods or food groups.³⁵
203 For example, the USDA database¹³ which was used to determine the vitamin K Nutrient
204 Reference Values for Australia and New Zealand.³⁷ Presently, the AI for vitamin K intake is
205 set at 60 and 70 µg/d for adult Australian females and males respectively.³⁷ Alternatively, it
206 has been proposed that even intakes of 90-120 µg/day (set by the USDA)¹³ of vitamin K may
207 not be sufficient to induce complete carboxylation of all VKDPs.³⁸ Consequently, AI
208 recommendations in Australia for vitamin K intake may not accurately reflect actual vitamin
209 K intake (or demands), and may lead to ambiguity when determining optimal vitamin K
210 nutrition. There is a need for both the development of regional databases and the expansion of
211 existing databases to include both vitamin K1 and K2 content (including a range of MKs) in
212 commonly consumed foods. Such work will allow for improvements in the determination of
213 dietary vitamin K recommendations. Finally, due to differences in the bioavailability and/or

214 bioaccessibility of vitamin K2, future guidelines may need to consider distinguishing between
215 dietary vitamin K1 and K2.

216 *Vitamin K databases*

217 Databases of nutrients allow for the calculation of individual and population level intake of
218 nutrients, permitting for epidemiological investigation. Although there appears to be ~70
219 sources containing information relating to the vitamin K content of food, closer analysis of
220 these reveal that current databases are largely incomplete.²⁵ Specifically, only 12 databases list
221 the vitamin K content of individual food items, which is required to more accurately determine
222 vitamin K intake.²⁵ The Dutch database is the most comprehensive and includes PK and several
223 types of MKs, ranging from MK-4 to MK-10.³⁹ Although the largest database (in English) has
224 been developed by the USDA, this database does have limitations. Specifically, food items
225 have only been assessed for PK and MK-4 in the 25th release of the USDA database.¹³
226 Considering no other MKs have been included in the database, this limits its utility for
227 accurately estimating total vitamin K2 intake. The USDA database also lacks some commonly
228 consumed animal products, including lamb, veal and goat, which are potentially rich sources
229 of MKs.¹³ The limited availability of comprehensive, national databases that lists both PK and
230 MK content in food hampers investigation of the relationship between dietary vitamin K1
231 and/or K2 intake and a range of health outcomes. Nevertheless, available databases provide
232 researchers with an indication of the types of food that are rich in vitamin K1 and K2 (including
233 the specific MKs). By reviewing pre-existing databases,^{3, 13, 15, 17, 39, 40} we have compiled a list
234 of commonly consumed foods and their approximate vitamin K content in *Table 1*. However,
235 estimates of the respective vitamin K1 and K2 content of the foods listed in *Table 1* should be
236 interpreted with caution, as there is substantial variability of PK and MK content in food by
237 region.²

238

239 *Regional differences in the vitamin K content of food*

240 The content of both PK and MKs in foods are known to differ between regions.² For example,
241 beef cuts in Japan contain more MK-4 than the United States (15.0 ± 7.0 vs. 1.1 to 9.3 $\mu\text{g}/100$
242 g, respectfully).^{14, 15} Similarly, the MK-4 content in egg yolk is four times higher in Japan
243 compared to the United States (64 vs 15.5 $\mu\text{g}/100\text{g}$).² The variability in MK-4 has been
244 attributed to the use of menadione in animal feeds and further differences in food production.²
245 Variability of up to 15% has been reported in the MK content (MK-6 to MK-10) of semi-hard
246 cheese varieties in three different European countries (France, Poland, Denmark).¹⁶
247 Furthermore, intra-country MK variability can be high (up to 80%), as reported between six
248 English cheddar cheeses.¹⁶ Results are also complicated by a range of methodologies
249 (discussed in subsequent section) used to quantify PK and MKs. The regional differences in
250 food production, climate conditions, and dietary consumption patterns highlights the
251 importance of comprehensive and detailed databases that are population specific. Ideally,
252 databases should include information relating to the brand, type, method of creation and region
253 of grown foods alongside their PK and MK content. If regional differences in vitamin K content
254 of food are ignored, this can lead to inaccuracies when estimating vitamin K intake. Although
255 the investment in comprehensive databases is unlikely to occur in most countries, the creation
256 of more region specific databases on the vitamin K contents of foods would ultimately facilitate
257 better estimation of vitamin K1 and K2 intake.

258 **Measuring dietary vitamin K**

259 *Methods to quantitate phylloquinone and menaquinones in food*

260 To develop databases, a validated method of measuring PK and MKs simultaneously, with
261 accuracy and high throughput, is required. These methods must have high sensitivity and
262 selectivity due to the low vitamin K content in foods, and the complexity of the food matrix.⁴¹
263 A recent review concisely summarised quantification methods for determination of vitamin K

264 in various biological matrices including blood, urine and tissue.⁴² However, the quantification
265 of vitamin K from food were not considered in detail. Existing methods to determine PK and
266 MK levels from food generally use reverse-phase HPLC with fluorescent detection or HPLC–
267 Mass Spectrometry (MS), with K1 or deuterated-K1 as the internal standard, respectively.^{16, 43}
268 Gas Chromatography-MS (GC-MS) has also been used with deuterium-labelled internal
269 standards to accurately measure PK in serum.^{44, 45} However, there are limitations in using GC
270 for measuring vitamin K, as high temperatures are needed to volatise vitamin K compounds
271 (>300°C).⁴⁶ Therefore, HPLC is the preferred method to separate vitamin K compounds for
272 analysis.⁴¹ The emerging method to measure vitamin K in foods is a HPLC-MS method
273 developed by Karl et al.⁴³ This method has high versatility and may be used to measure PK and
274 all MKs in a variety of forms such as food, serum and faeces.⁴³ Based on current evidence,
275 HPLC-MS appears to be the best method to measure vitamin K compounds from foods.

276 *Methods of extracting vitamin K from the food matrix*

277 Vitamin K compounds are trapped in the matrix of foods and must be liberated in order to be
278 accurately measured.³ Vitamin K compounds are unstable in alkali conditions and undergo
279 photo-oxidation upon exposure to ultraviolet light, which limits the available extraction
280 methods.⁴⁶ Samples must be stored in amber vials and all work done yellow light to prevent
281 this degradation.⁴³ Due to these limitations, the most common extraction method is liquid-
282 liquid extraction, with the exact process depending on the food matrix.^{46, 47} Sample pre-
283 treatment is important to liberate K vitamers from the matrix. Most food groups, with the
284 exception of vegetable oils, are homogenised and sonicated to break up the food matrix.^{3, 48}
285 This is especially important for vegetables, as it is essential to liberate PK from the thylakoid
286 membranes.⁴⁶ Furthermore, some fatty foods, such as oils or cheeses, undergo lipase digestion
287 or acid treatment to break down fats, thereby increasing purity.^{16, 46} To our knowledge, there is

288 currently no standardised method for the extraction of vitamin K compounds from food,
289 however, numerous validated methods exist.^{3, 43, 47}

290 **Vitamin K and cardiovascular health**

291 *Atherosclerotic cardiovascular disease*

292 Cardiovascular disease accounts for approximately 1 in 3 deaths worldwide.⁴⁹ Atherosclerotic
293 cardiovascular disease includes coronary heart disease (CHD), peripheral arterial disease and
294 cerebrovascular disease. Atherosclerotic diseases are caused by endothelial dysfunction,
295 inflammation and the slow build-up of cholesterol, fats and calcium deposits forming plaques
296 in arterial walls.⁵⁰ The plaques decrease lumen area of the artery, increasing resistance to blood
297 flow. Vulnerable plaques can rupture or erode leading to blood clots, which can obstruct blood
298 flow to the heart causing a heart attack, or to the brain causing a stroke. These plaques frequently
299 develop calcium mineral deposits, which further adds to plaque volume and leads to reduced
300 arterial compliance leading to an increased risk of organ damage.⁵⁰ As such, strategies capable
301 of preventing or delaying the progression of atherosclerosis are fundamental to improving
302 cardiovascular health.

303 *Vitamin K and vascular calcification*

304 The arterial (vascular) calcification process was once thought to be passive and a natural
305 consequence of ageing. However, it is now understood to be a complex biomineralisation
306 process which is actively regulated.⁵¹ Observational studies have reported that vitamin K intake
307 is inversely associated with arterial calcification.⁵² The best evidence for a causal role for
308 vitamin K in arterial calcification comes from vitamin K antagonists (VKA), genetic studies
309 and short-term randomised controlled trials.⁵²⁻⁵⁴ VKA such as warfarin are anticoagulants used
310 to prevent thrombosis and pulmonary emboli. VKAs reduce the bioactivation of the hepatic
311 VKDP, such as prothrombin, which contributes to coagulation.⁵⁵ However, their effects are
312 non-specific and systemic, thus extrahepatic VKDP are inactivated during VKA treatment.^{54,}

313 ⁵⁶ This includes Matrix-Gla Protein (MGP), an inhibitor of arterial calcification,⁵⁷ activated
314 protein-C an inhibitor of inflammation and endothelial cell apoptosis,⁵⁸ and osteocalcin that
315 may regulate metabolic dysfunction and arterial calcification.⁵⁹ Studies have shown that the
316 long-term use of VKA is associated with accelerated arterial calcification.^{2, 60} VKDPs have
317 high affinity for calcium-based matrices through their Gla residues⁵⁸ and the potential role of
318 vitamin K in enhancing/activating anti-calcification processes (especially through MGP) has
319 gained much interest.¹ However, a fine balance may exist as anti-thrombotic blocking agents
320 of vitamin K on blood coagulation are an important treatment for preventing heart attacks and
321 strokes. Although controversial, this has led to suggestions that individuals on anti-coagulants
322 should consider avoiding vitamin K rich foods.^{61, 62}

323 MGPs ability to reduce arterial calcification has been demonstrated in MGP knockout mice,
324 which rapidly develop severe calcification and die from blood-vessel rupture.⁶³ Additionally,
325 patients suffering from Keutel syndrome, where mutations to the MGP gene render the protein
326 non-functional,⁶⁴ also develop extensive soft tissue calcification.⁶⁵ Polymorphisms within the
327 MGP have been associated with both atherosclerosis and vascular calcification.^{53, 66} As
328 summarised in *Figure 2*, these results suggest that VKDP, and in particular MGP, may play a
329 pivotal role in preventing vascular calcification in humans. As a result of the aforementioned
330 work, the importance of vitamin K for cardiovascular health has recently been investigated in
331 several observational studies and clinical trials.

332 *Observational studies of vitamin K intake and clinical cardiovascular disease*

333 To date, there are few prospective studies investigating the role of dietary vitamin K intake on
334 the risk of CVD. These studies are particularly challenging as most databases have very limited
335 data on the MK content in food, presenting a significant limitation. Likewise, PK intake is
336 susceptible to social bias, because vegetable intake is generally over-reported, and is likely to
337 be influenced by variation in vegetable intake across populations.⁶⁷ Validating the FFQs used

338 to estimate vitamin K2 intake is also challenging, as MKs are not typically detected in the
339 circulation unless large quantities of MK-rich foods are consumed.⁶⁸ Furthermore, dietary PK
340 is likely to be influenced by variation in vegetable intake across populations. The Rotterdam
341 Study (n= 4807), one of the largest investigations to date, found that the relative risk (RR) of
342 CHD mortality was reduced for individuals within the middle (21.6–32.7 µg/d) and upper
343 (>32.7 µg/d) tertiles (RR 0.73 95%CI: 0.45-1.17 and RR 0.43 95%CI 0.24-0.77, respectively)
344 compared to the lowest tertile (<21.6 µg/d) of vitamin K2 intake. Similarly, individuals in the
345 highest tertile of vitamin K2 intake had lower odds (OR 0.48 95%CI 0.32-0.71) for severe
346 aortic calcification compared to individuals in the lowest tertile of vitamin K2 intake.⁶⁹ Of
347 interest, in the Rotterdam study vitamin K1 intake was not related to CHD. However, this was
348 based on 233 events and there was a statistically non-significant 11% lower relative hazard
349 (HR 0.89 95%CI 0.63-1.25) for individuals with the highest intake of PK.⁶⁹ Noteworthy, this
350 data should be interpreted with caution as the reproducibility and validity of the FFQ used in
351 the Rotterdam study for estimating vitamin K intake has been reported to be problematic.⁷⁰ The
352 finding that higher dietary K2 is inversely related (per 10 µg/d, HR 0.91 95%CI 0.85-1.00) to
353 incident CHD has been replicated in the Prospect-EPIC cohort of 16,057 women (aged 49–70
354 years) free of cardiovascular diseases at baseline.⁷¹ This protective affect was attributed
355 predominantly to long-chain vitamin K2 subtypes MK-7, MK-8 and MK-9.⁷¹

356 Higher dietary MK intake (per 10 µg/d) has also been associated with an 8% decrease in the
357 relative hazard (HR 0.92 95%CI 0.85-0.99) for peripheral arterial disease (PAD) in 36,629
358 Dutch individuals from the Prospect-EPIC and MORGEN-EPIC cohorts.⁷² In contrast, some
359 studies have reported no association between vitamin K2 intake and CVD mortality or stroke.⁷³⁻
360 ⁷⁵ It should be noted that the majority of these observational studies have been performed in
361 the Netherlands; likely due to the Dutch having one of the most comprehensive databases for
362 vitamin K (especially K2) content in food.⁶⁹ In contrast to MKs, high PK intake has not been

363 associated with CVD mortality,⁷³ CHD incidence and mortality^{69, 71, 73, 76, 77} and PAD⁷².
364 However, in two studies where inverse associations between PK intake and CHD incidence
365 were identified, this relationship was attenuated once other dietary factors were considered.^{76,}
366 ⁷⁷ This suggests that other components of the diet may also be contributing to observed
367 findings, and that the healthy dietary patterns associated with high PK intake may at least
368 partially explain any relationship with reduced CHD risk, rather than PK itself.^{76,77} Similarly,
369 studies examining the link between MK and CHD may be influenced by other components in
370 MK-rich foods (e.g. saturated fats and sodium from processed meats) associated with CHD.
371 Additionally, the lack of comprehensive databases for vitamin K1 and K2, and more
372 specifically the long-chain MK (e.g. MK-5 to MK-13) content of commonly consumed foods
373 limits our understanding of the role of vitamin K on vascular ageing and clinical cardiovascular
374 events. As such, caution needs to be exercised when interpreting epidemiological studies of
375 vitamin K with CHD outcomes.

376 *Vitamin K supplementation trials with surrogate vascular outcomes*

377 To our knowledge, no RCT have examined the effects of vitamin K (e.g. non-pharmaceutical)
378 on clinical CVD outcomes. Here we discuss the evidence from RCTs investigating surrogate
379 vascular outcomes such as coronary artery calcification (CAC), arterial stiffness and aortic
380 valve calcification (AVC). A short 8-week open-label single-arm trial in renal transplant
381 patients (n= 60) reported a significant decrease in arterial stiffness and 14.2% reduction in
382 mean carotid femoral pulse wave velocity with higher doses (360 µg/day) of MK-7
383 supplementation. Of interest, the most benefit was gained in patients presenting with a vitamin
384 K deficiency (undercarboxylated-MGP >500 pmol/L) at baseline.⁷⁸ Similar findings were
385 reported in a separate single arm study that supplemented individuals (n=26, with at least one
386 coronary risk factor) with 45 µg/day of MK-4 for one year. Specifically, reduced arterial
387 stiffness (but no changes on CAC) were recorded only in the subset of vitamin K deficient

388 patients (n=4).⁷⁹ The aforementioned work indicate that the vitamin K status of populations
389 need to be considered when examining potential beneficial of vitamin K for vascular outcomes.
390 A double blind, placebo-controlled trial found MK-7 supplementation (180 µg/d) over three
391 years reduced arterial stiffness in 244 healthy post-menopausal women. Furthermore, in
392 women with higher arterial stiffness (stiffness index β scores ≥ 10.8) at baseline, improvements
393 in local carotid pulse wave velocity were recorded, suggesting better elasticity.⁸⁰ Contrary to
394 the aforementioned positive findings, a recent double blind, placebo-controlled trial
395 supplementing MK-7 (360 µg/d) over 6 months in older individuals (~69 y, 24% female) with
396 type II diabetes and CVD found no significant reduction in femoral arterial calcification
397 measured by sodium fluoride positron emission tomography.⁸¹

398 To date, few trials have used PK supplements and tracked cardiovascular outcomes.⁷² An open-
399 label randomised placebo controlled trial of 72 (82% male, mean age 69 years) individuals
400 reported that 12 months of high dose PK supplementation (2 mg/d) in patients with
401 asymptomatic or mildly symptomatic aortic valve calcification (AVC) reduced AVC
402 progression by ~12%.⁸² Another 3-year double-blind placebo controlled trial examined the
403 effect of a multivitamin supplement with 500 µg/d of PK (n=200) vs. an identical multivitamin
404 supplement without PK (n=188) provided in healthy post-menopausal women and men.⁸³ In
405 participants with high adherence to the supplements ($\geq 85\%$, n=367), there was less CAC
406 progression in the PK group than in the control group. Furthermore, in individuals with pre-
407 existing CAC (Agatston score >10), those supplemented with PK had 6% less CAC
408 progression. A randomised placebo controlled trial in postmenopausal women (n=150)
409 receiving a supplement containing PK (1 mg/day) and vitamin D for 3 years demonstrated
410 better carotid artery compliance and elasticity, measured by ultrasound of the common carotid
411 artery.⁸⁴ Specifically, when comparing the vitamin K and D supplemented group to the placebo
412 group, better arterial compliance coefficient (8.6%), distensibility coefficient (8.8%), and pulse

413 pressure (6.3%) were recorded. Collectively, these large population studies and RCTs suggest
414 that vitamin K (PK and/or MKs) supplementation may play a vital role in blood vessel health.
415 An important consideration for future work is to examine if the aforementioned cardiovascular
416 health benefits observed after PK and/or MK supplementation may be limited only to
417 populations presenting with vitamin K insufficiency and/or deficiency.

418 **Conclusion**

419 Until recently, a large proportion of research into vitamin K often only considered PK, with
420 the role of MKs remaining largely ambiguous. It is still unclear as to what the most appropriate
421 dietary recommendations for daily vitamin K should be, including if a differentiation between
422 vitamin K1 and K2 need to be considered in nutritional guidelines promoted by public health
423 organisations. Due to difficulties in the measurement of vitamin K2, the MK derivatives have
424 also not been extensively tested for basic pharmacokinetic data or their concentration in
425 numerous food items. Presently, epidemiological data suggest that high dietary MK intake may
426 be protective against CHD mortality and coronary artery calcification.⁶⁹ However, the bulk of
427 evidence for health benefits are limited to the Netherlands; as it currently possesses the most
428 comprehensive vitamin K database, particularly for MKs. Therefore, observational studies are
429 still needed to examine these findings in countries with different dietary patterns. There is also
430 limited evidence from RCTs highlighting the benefit of PK supplements in regards to delaying
431 the progression of CAC, and improving carotid artery compliance.^{52, 84} Further investigation
432 using vitamin K supplementation and its effect on clinical outcomes are still needed.
433 Unfortunately, current research relating to vitamin K is hampered by the absence of
434 comprehensive national databases that list both PK and MK content in food. However, with
435 recent advances in the measurement of PK and MKs using HPLC-MS,⁴³ this could lead to the
436 development of comprehensive region-specific vitamin K databases. This is especially
437 important as regional variability in vitamin K content in food may limit the use of out of region

438 vitamin K databases.² Such work will also enable future investigations to explore the potential
439 importance of dietary PK and MKs (separately or collectively) intake for a range of health
440 outcomes. The application of these databases to large cohort studies will advance our
441 understanding of the importance of vitamin K for human health, especially in the
442 cardiovascular field. This will inform well design RCTs to establish causal effects between
443 dietary vitamin K and a range of health outcomes.

444 **Conflicts of interest**

445 There are no conflicts of interest to declare.

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451 **References**

- 452 1. J. J. DiNicolantonio, J. Bhutani and J. H. O'Keefe, The health benefits of vitamin K,
453 *Open Heart*, 2015, **2**, DOI: 10.1136/openhrt-2015-000300.
- 454 2. B. Walther, J. P. Karl, S. L. Booth and P. Boyaval, Menaquinones, bacteria, and the
455 food supply: the relevance of dairy and fermented food products to vitamin K
456 requirements, *Adv. Nutr.*, 2013, **4**, 463-473.
- 457 3. L. J. Schurgers and C. Vermeer, Determination of phylloquinone and menaquinones
458 in food, *Pathophysiol. Haemost. Thromb.*, 2000, **30**, 298-307.
- 459 4. M. J. Shearer and P. Newman, Recent trends in the metabolism and cell biology of
460 vitamin K with special reference to vitamin K cycling and MK-4 biosynthesis, *J.*
461 *Lipid Res.*, 2014, **55**, 345-362.
- 462 5. B. Nowicka and J. Kruk, Occurrence, biosynthesis and function of isoprenoid
463 quinones, *Biochim. Biophys. Acta*, 2010, **1797**, 1587-1605.
- 464 6. G. Lippi and M. Franchini, Vitamin K in neonates: facts and myths, *Blood Transfus.*,
465 2011, **9**, 4-9.
- 466 7. C. W. Thane, A. A. Paul, C. J. Bates, C. Bolton-Smith, A. Prentice and M. J. Shearer,
467 Intake and sources of phylloquinone (vitamin K1): variation with socio-demographic
468 and lifestyle factors in a national sample of british elderly people, *Br. J. Nutr.*, 2002,
469 **87**, 605-613.
- 470 8. C. Bolton-Smith, R. J. G. Price, S. T. Fenton, D. J. Harrington and M. J. Shearer,
471 Compilation of a provisional UK database for the phylloquinone (vitamin K1) content
472 of foods, *Br. J. Nutr.*, 2007, **83**, 389-399.
- 473 9. S. G. Harshman, E. G. Finnan, K. J. Barger, R. L. Bailey, D. B. Haytowitz, C. H.
474 Gilhooly and S. L. Booth, Vegetables and mixed dishes are top contributors to
475 phylloquinone intake in US adults: Data from the 2011-2012 NHANES, *J. Nutr.*,
476 2017, **147**, 1308-1313.
- 477 10. K. Nimptsch, S. Rohrmann and J. Linseisen, Dietary intake of vitamin K and risk of
478 prostate cancer in the heidelberg cohort of the european prospective investigation into
479 cancer and nutrition (EPIC-Heidelberg), *Am. J. Clin. Nutr.*, 2008, **87**, 985-992.
- 480 11. L. J. Schurgers, J. M. Geleijnse, D. E. Grobbee, H. A. P. Pols, A. Hofman, J. C. M.
481 Witteman and C. Vermeer, Nutritional intake of vitamins K1 (phylloquinone) and K2
482 (menaquinone) in the netherlands, *J. Nutr. Environ. Med.*, 1999, **9**, 115-122.

- 483 12. X. Fu, X. Shen, E. G. Finnan, D. B. Haytowitz and S. L. Booth, Measurement of
484 multiple vitamin K forms in processed and fresh-cut pork products in the U.S. food
485 supply, *J. Agric. Food Chem.*, 2016, **64**, 4531-4535.
- 486 13. USDA provisional table on the vitamin K contents in food,
487 [https://ndb.nal.usda.gov/ndb/nutrients/report/nutrientsfrm?max=25&offset=0&totCou](https://ndb.nal.usda.gov/ndb/nutrients/report/nutrientsfrm?max=25&offset=0&totalCount=0&nutrient1=430&nutrient2=428&nutrient3=&subset=0&sort=c&measureby=g)
488 [nt=0&nutrient1=430&nutrient2=428&nutrient3=&subset=0&sort=c&measureby=g,](https://ndb.nal.usda.gov/ndb/nutrients/report/nutrientsfrm?max=25&offset=0&totalCount=0&nutrient1=430&nutrient2=428&nutrient3=&subset=0&sort=c&measureby=g)
489 (accessed March 30, 2019).
- 490 14. S. J. Elder, D. B. Haytowitz, J. Howe, J. W. Peterson and S. L. Booth, Vitamin K
491 contents of meat, dairy, and fast food in the U.S. diet, *J. Agric. Food Chem.*, 2006, **54**,
492 463-467.
- 493 15. M. Kamao, Y. Suhara, N. Tsugawa, M. Uwano, N. Yamaguchi, K. Uenishi, H. Ishida,
494 S. Sasaki and T. Okano, Vitamin K content of foods and dietary vitamin K intake in
495 Japanese young women, *J. Nutr. Sci. Vitaminol. (Tokyo)*, 2007, **53**, 464-470.
- 496 16. E. Manoury, K. Jourdon, P. Boyaval and P. Fourcassie, Quantitative measurement of
497 vitamin K2 (menaquinones) in various fermented dairy products using a reliable high-
498 performance liquid chromatography method, *J. Dairy Sci.*, 2013, **96**, 1335-1346.
- 499 17. X. Fu, S. G. Harshman, X. Shen, D. B. Haytowitz, J. P. Karl, B. E. Wolfe and S. L.
500 Booth, Multiple vitamin K forms exist in dairy foods, *Curr Dev Nutr*, 2017, **1**, DOI:
501 10.3945/cdn.117.000638.
- 502 18. J. W. J. Beulens, M. L. Bots, F. Atsma, M.-L. E. L. Bartelink, M. Prokop, J. M.
503 Geleijnse, J. C. M. Witteman, D. E. Grobbee and Y. T. van Der Schouw, High dietary
504 menaquinone intake is associated with reduced coronary calcification,
505 *Atherosclerosis*, 2009, **203**, 489-493.
- 506 19. B. Holst and G. Williamson, Nutrients and phytochemicals: from bioavailability to
507 bioefficacy beyond antioxidants, *Curr. Opin. Biotechnol.*, 2008, **19**, 73-82.
- 508 20. J. Beulens, S. Booth, E. van Den Heuvel, E. Stoecklin, A. Baka and C. Vermeer, The
509 role of menaquinones (vitamin K2) in human health, *Br. J. Nutr.*, 2013, **110**, 1357-
510 1368.
- 511 21. S. L. Booth, A. H. Lichtenstein and G. E. Dallal, Phylloquinone absorption from
512 phylloquinone-fortified oil is greater than from a vegetable in younger and older men
513 and women, *J. Nutr.*, 2002, **132**, 2609-2612.
- 514 22. A. K. Garber, N. C. Binkley, D. C. Krueger and J. W. Suttie, Comparison of
515 phylloquinone bioavailability from food sources or a supplement in human subjects,
516 *J. Nutr.*, 1999, **129**, 1201-1203.

- 517 23. L. J. Schurgers and C. Vermeer, Differential lipoprotein transport pathways of K-
518 vitamins in healthy subjects, *Biochim. Biophys. Acta*, 2002, **1570**, 27-32.
- 519 24. T. Sato, L. J. Schurgers and K. Uenishi, Comparison of menaquinone-4 and
520 menaquinone-7 bioavailability in healthy women, *Nutr. J.*, 2012, **11**, 93.
- 521 25. M. C. Barbara Walther, in *Vitamin K2 - Vital for Health and Wellbeing*, ed. J. O.
522 Gordeladze, IntechOpen, 2017, DOI: 10.5772/63712.
- 523 26. K. Nakagawa, Y. Hirota, N. Sawada, N. Yuge, M. Watanabe, Y. Uchino, N. Okuda,
524 Y. Shimomura, Y. Suhara and T. Okano, Identification of UBIAD1 as a novel human
525 menaquinone-4 biosynthetic enzyme, *Nature*, 2010, **468**, 117-121.
- 526 27. Y. Hirota, N. Tsugawa, K. Nakagawa, Y. Suhara, K. Tanaka, Y. Uchino, A. Takeuchi,
527 N. Sawada, M. Kamao, A. Wada, T. Okitsu and T. Okano, Menadione (vitamin K3) is
528 a catabolic product of oral phylloquinone (vitamin K1) in the intestine and a
529 circulating precursor of tissue menaquinone-4 (vitamin K2) in rats, *J. Biol. Chem.*,
530 2013, **288**, 33071-33080.
- 531 28. M. J. Shearer, X. Fu and S. L. Booth, Vitamin K nutrition, metabolism, and
532 requirements: current concepts and future research, *Adv. Nutr.*, 2012, **3**, 182-195.
- 533 29. F. Fernandez and M. D. Collins, Vitamin K composition of anaerobic gut bacteria,
534 *FEMS Microbiol. Lett.*, 1987, **41**, 175-180.
- 535 30. M. D. Collins and D. Jones, Distribution of isoprenoid quinone structural types in
536 bacteria and their taxonomic implication, *Microbiol. Rev.*, 1981, **45**, 316-354.
- 537 31. M. J. Shearer and P. Newman, Metabolism and cell biology of vitamin K, *Thromb.*
538 *Haemost.*, 2008, **100**, 530-547.
- 539 32. J. M. Conly and K. Stein, Quantitative and qualitative measurements of K vitamins in
540 human intestinal contents, *Am. J. Gastroenterol.*, 1992, **87**, 311-316.
- 541 33. J. W. Suttie, The importance of menaquinones in human nutrition, *Annu. Rev. Nutr.*,
542 1995, **15**, 399-417.
- 543 34. H. K. Biesalski, Nutrition meets the microbiome: micronutrients and the microbiota,
544 *Ann. N. Y. Acad. Sci.*, 2016, **1372**, 53-64.
- 545 35. N. EFSA Panel on Dietetic Products, Allergies, D. Turck, J.-L. Bresson, B.
546 Burlingame, T. Dean, S. Fairweather-Tait, M. Heinonen, K. I. Hirsch-Ernst, I.
547 Mangelsdorf, H. J. McArdle, A. Naska, G. Nowicka, K. Pentieva, Y. Sanz, A. Siani,
548 A. Sjödin, M. Stern, D. Tomé, H. Van Loveren, M. Vinceti, P. Willatts, C. Lamberg-
549 Allardt, H. Przyrembel, I. Tetens, C. Dumas, L. Fabiani, S. Ioannidou and M.
550 Neuhäuser-Berthold, Dietary reference values for vitamin K, *EFSA J*, 2017, **15**, DOI:

- 551 10.2903/j.efsa.2017.4780.
- 552 36. *Nutrient reference values for Australia and New Zealand: including recommended*
553 *dietary intakes*, Report 1864962372, National Health and Medical Research Council,
554 Canberra, ACT, 2006.
- 555 37. M. Vetrella and W. Barthelmai, Studies on drug-induced hemolysis: Effects of
556 menadione and its water soluble preparations on the glutathione peroxidase of human
557 erythrocytes, *Klin. Wochenschr.*, 1972, **50**, 234-238.
- 558 38. N. C. Binkley, D. C. Krueger, T. N. Kawahara, J. A. Engelke, R. J. Chappell and J.
559 W. Suttie, A high phyloquinone intake is required to achieve maximal osteocalcin
560 gamma-carboxylation, *Am. J. Clin. Nutr.*, 2002, **76**, 1055-1060.
- 561 39. S. Westenbrink and M. Jansen-van der Vliet, Dutch Food Composition Table,
562 <https://www.rivm.nl/en/dutch-food-composition-database>, (accessed March 30, 2019).
- 563 40. C. Vermeer, J. Raes, C. van 't Hoofd, M. H. J. Knapen and S. Xanthoulea,
564 Menaquinone content of cheese, *Nutrients*, 2018, **10**, DOI: 10.3390/nu10040446.
- 565 41. Y. Zhang, W.-E. Zhou, J.-Q. Yan, M. Liu, Y. Zhou, X. Shen, Y.-L. Ma, X.-S. Feng, J.
566 Yang and G.-H. Li, A Review of the extraction and determination methods of thirteen
567 essential vitamins to the human body: An update from 2010, *Molecules*, 2018, **23**,
568 DOI: 10.3390/molecules23061484.
- 569 42. Y. Zhang, V. Bala, Z. Mao, Y. S. Chhonker and D. J. Murry, A concise review of
570 quantification methods for determination of vitamin K in various biological matrices,
571 *J. Pharm. Biomed. Anal.*, 2019, **169**, 133-141.
- 572 43. J. P. Karl, X. Fu, G. G. Dolnikowski, E. Saltzman and S. L. Booth, Quantification of
573 phyloquinone and menaquinones in feces, serum, and food by high-performance
574 liquid chromatography-mass spectrometry, *J. Chromatogr. B Analyt. Technol.*
575 *Biomed. Life Sci.*, 2014, **963**, 128-133.
- 576 44. G. G. Dolnikowski, Z. Sun, M. A. Grusak, J. W. Peterson and S. L. Booth, HPLC and
577 GC/MS determination of deuterated vitamin K (phyloquinone) in human serum after
578 ingestion of deuterium-labeled broccoli, *J. Nutr. Biochem.*, 2002, **13**, 168-174.
- 579 45. K. S. Jones, L. J. Bluck and W. A. Coward, Analysis of isotope ratios in vitamin K1
580 (phyloquinone) from human plasma by gas chromatography/mass spectrometry,
581 *Rapid Commun. Mass Spectrom.*, 2006, **20**, 1894-1898.
- 582 46. T. Koivu-Tikkanen, Determination of phyloquinone and menaquinones in foods by
583 HPLC, University of Helsinki, 2001.

- 584 47. S. L. Booth and J. A. Sadowski, in *Methods in Enzymology*, Academic Press, 1997,
585 Determination of phylloquinone in foods by high-performance liquid
586 chromatography, 446-456.
- 587 48. A. Gentili, A. Cafolla, T. Gasperi, S. Bellante, F. Caretti, R. Curini and V. P.
588 Fernandez, Rapid, high performance method for the determination of vitamin K1,
589 menaquinone-4 and vitamin K1 2,3-epoxide in human serum and plasma using liquid
590 chromatography-hybrid quadrupole linear ion trap mass spectrometry, *J. Chromatogr.*
591 *A*, 2014, **1338**, 102-110.
- 592 49. World Health Organisation, Cardiovascular diseases fact sheet, 2017.
- 593 50. S. Mendis, P. Puska and B. Norrving, *Global atlas on cardiovascular disease*
594 *prevention and control*, World Health Organisation, 2011.
- 595 51. M. Abedin, Y. Tintut and L. Demer Linda, Vascular calcification, *Arterioscler.*
596 *Thromb. Vasc. Biol.*, 2004, **24**, 1161-1170.
- 597 52. M. K. Shea and R. M. Holden, Vitamin K status and vascular calcification: evidence
598 from observational and clinical studies, *Adv. Nutr.*, 2012, **3**, 158-165.
- 599 53. K. Sheng, P. Zhang, W. Lin, J. Cheng, J. Li and J. Chen, Association of matrix gla
600 protein gene (rs1800801, rs1800802, rs4236) polymorphism with vascular
601 calcification and atherosclerotic disease: a meta-analysis, *Sci. Rep.*, 2017, **7**, DOI:
602 10.1038/s41598-017-09328-5.
- 603 54. B. A. Willems, C. Vermeer, C. P. Reutelingsperger and L. J. Schurgers, The realm of
604 vitamin K dependent proteins: shifting from coagulation toward calcification, *Mol.*
605 *Nutr. Food Res.*, 2014, **58**, 1620-1635.
- 606 55. J. Danziger, Vitamin K-dependent proteins, warfarin, and vascular calcification, *Clin.*
607 *J. Am. Soc. Nephrol.*, 2008, **3**, 1504-1510.
- 608 56. E. Theuwissen, E. Smit and C. Vermeer, The role of vitamin K in soft-tissue
609 calcification, *Adv. Nutr.*, 2012, **3**, 166-173.
- 610 57. H. Barrett, M. O'Keeffe, E. Kavanagh, M. Walsh and E. M. O'Connor, Is matrix gla
611 protein associated with vascular calcification? a systematic review, *Nutrients*, 2018,
612 **10**, DOI: 10.3390/nu10040415.
- 613 58. J. H. Griffin, J. A. FernÁNdez, A. J. Gale and L. O. Mosnier, Activated protein C, *J.*
614 *Thromb. Haemost.*, 2007, **5**, 73-80.
- 615 59. S. L. Booth, A. Centi, S. R. Smith and C. Gundberg, The role of osteocalcin in human
616 glucose metabolism: marker or mediator?, *Nat. Rev. Endocrinol.*, 2013, **9**, 43-55.

- 617 60. L. J. Schurgers, H. Aebert, C. Vermeer, B. Bültmann and J. Janzen, Oral
618 anticoagulant treatment: friend or foe in cardiovascular disease?, *Blood*, 2004, **104**,
619 3231-3232.
- 620 61. F. Violi, G. Y. Lip, P. Pignatelli and D. Pastori, Interaction between dietary vitamin K
621 intake and anticoagulation by vitamin K antagonists: is it really true?: A systematic
622 review, *Medicine (Baltimore)*, 2016, **95**, DOI: 10.1097/MD.0000000000002895.
- 623 62. C. M. Schooling, Plasma levels of vitamin K and the risk of ischemic heart disease: a
624 mendelian randomization study, *J. Thromb. Haemost.*, 2016, **14**, 1211-1215.
- 625 63. G. Luo, P. Ducy, M. D. McKee, G. J. Pinero, E. Loyer, R. R. Behringer and G.
626 Karsenty, Spontaneous calcification of arteries and cartilage in mice lacking matrix
627 GLA protein, *Nature*, 1997, **386**, 78-81.
- 628 64. P. B. Munroe, R. O. Olgunturk, J.-P. Fryns, L. V. Maldergem, F. Ziereisen, B.
629 Yuksel, R. M. Gardiner and E. Chung, Mutations in the gene encoding the human
630 matrix gla protein cause keutel syndrome, *Nat. Genet.*, 1999, **21**, 142-144.
- 631 65. M. Meier, L. P. Weng, E. Alexandrakis, J. Rüschoff and G. Goeckenjan,
632 Tracheobronchial stenosis in keutel syndrome, *Eur. Respir. J.*, 2001, **17**, 566-569.
- 633 66. S. M. Herrmann, C. Whatling, E. Brand, V. Nicaud, J. Gariépy, A. Simon, A. Evans,
634 J. B. Ruidavets, D. Arveiler, G. Luc, L. Tiret, A. Henney and F. Cambien,
635 Polymorphisms of the human matrix gla protein (MGP) gene, vascular calcification,
636 and myocardial infarction, *Arterioscler. Thromb. Vasc. Biol.*, 2000, **20**, 2386-2393.
- 637 67. T. M. Miller, M. F. Abdel-Maksoud, L. A. Crane, A. C. Marcus and T. E. Byers,
638 Effects of social approval bias on self-reported fruit and vegetable consumption: a
639 randomized controlled trial, *Nutr. J.*, 2008, **7**, DOI: 10.1186/1475-2891-7-18.
- 640 68. M. K. Shea and S. L. Booth, Concepts and controversies in evaluating vitamin K
641 status in population-based studies, *Nutrients*, 2016, **8**, DOI: 10.3390/nu8010008.
- 642 69. J. M. Geleijnse, C. Vermeer, D. E. Grobbee, L. J. Schurgers, M. H. Knapen, I. M. van
643 der Meer, A. Hofman and J. C. Witteman, Dietary intake of menaquinone is
644 associated with a reduced risk of coronary heart disease: the rotterdam Study, *J. Nutr.*,
645 2004, **134**, 3100-3105.
- 646 70. S. R. Zwakenberg, A. I. P. Engelen, G. W. Dalmeijer, S. L. Booth, C. Vermeer, J.
647 Drijvers, M. C. Ocke, E. J. M. Feskens, Y. T. van der Schouw and J. W. J. Beulens,
648 Reproducibility and relative validity of a food frequency questionnaire to estimate
649 intake of dietary phylloquinone and menaquinones, *Eur. J. Clin. Nutr.*, 2017, **71**,
650 1423-1428.

- 651 71. G. C. M. Gast, N. M. de Roos, I. Sluijs, M. L. Bots, J. W. J. Beulens, J. M. Geleijnse,
652 J. C. Witteman, D. E. Grobbee, P. H. M. Peeters and Y. T. van der Schouw, A high
653 menaquinone intake reduces the incidence of coronary heart disease, *Nutr., Metab.*
654 *Cardiovasc. Dis.*, 2009, **19**, 504-510.
- 655 72. L. E. T. Vissers, G. W. Dalmeijer, J. M. A. Boer, W. M. M. Verschuren, Y. T. van
656 Der Schouw and J. W. J. Beulens, The relationship between vitamin K and peripheral
657 arterial disease, *Atherosclerosis*, 2016, **252**, 15-20.
- 658 73. S. R. Zwakenberg, N. R. den Braver, A. I. P. Engelen, E. J. M. Feskens, C. Vermeer,
659 J. M. A. Boer, W. M. M. Verschuren, Y. T. van der Schouw and J. W. J. Beulens,
660 Vitamin K intake and all-cause and cause specific mortality, *Clin. Nutr.*, 2017, **36**,
661 1294-1300.
- 662 74. L. E. T. Vissers, G. W. Dalmeijer, J. M. A. Boer, W. M. Monique Verschuren, Y. T.
663 van Der Schouw and J. W. J. Beulens, Intake of dietary phylloquinone and
664 menaquinones and risk of stroke, *J. Am. Heart Assoc.*, 2013, **2**, e000455.
- 665 75. M. Juanola-Falgarona, J. Salas-Salvado, M. A. Martinez-Gonzalez, D. Corella, R.
666 Estruch, E. Ros, M. Fito, F. Aros, E. Gomez-Gracia, M. Fiol, J. Lapetra, J. Basora, R.
667 M. Lamuela-Raventos, L. Serra-Majem, X. Pinto, M. A. Munoz, V. Ruiz-Gutierrez, J.
668 Fernandez-Ballart and M. Bullo, Dietary intake of vitamin K is inversely associated
669 with mortality risk, *J. Nutr.*, 2014, **144**, 743-750.
- 670 76. A. T. Erkkilä, S. L. Booth, F. B. Hu, P. F. Jacques, J. E. Manson, K. M. Rexrode, M.
671 J. Stampfer and A. H. Lichtenstein, Phylloquinone intake as a marker for coronary
672 heart disease risk but not stroke in women, *Eur. J. Clin. Nutr.*, 2004, **59**, 196.
- 673 77. A. T. Erkkilä, S. L. Booth, F. B. Hu, P. F. Jacques and A. H. Lichtenstein,
674 Phylloquinone intake and risk of cardiovascular diseases in men, *Nutr., Metab.*
675 *Cardiovasc. Dis.*, 2007, **17**, 58-62.
- 676 78. A. G. Mansour, E. Hariri, Y. Daaboul, S. Korjian, A. El Alam, A. D. Protogerou, H.
677 Kilany, A. Karam, A. Stephan and S. A. Bahous, Vitamin K2 supplementation and
678 arterial stiffness among renal transplant recipients—a single-arm, single-center
679 clinical trial, *J. Am. Soc. Hypertens.*, 2017, **11**, 589-597.
- 680 79. Y. Ikari, S. Torii, A. Shioi and T. Okano, Impact of menaquinone-4 supplementation
681 on coronary artery calcification and arterial stiffness: an open label single arm study,
682 *Nutr. J.*, 2016, **15**, DOI: 10.1186/s12937-016-0175-8.
- 683 80. M. H. J. Knapen, L. A. J. L. M. Braam, N. E. Drummen, O. Bekers, A. P. G. Hoeks
684 and C. Vermeer, Menaquinone-7 supplementation improves arterial stiffness in

- 685 healthy postmenopausal women. A double-blind randomised clinical trial, *Thromb.*
686 *Haemost.*, 2015, **113**, 1135-1144.
- 687 81. S. R. Zwakenberg, P. A. de Jong, J. W. Bartstra, R. van Asperen, J. Westerink, H.
688 de Valk, R. H. J. A. Slart, G. Luurtsema, J. M. Wolterink, G. J. de Borst, J. A.
689 van Herwaarden, M. A. van de Ree, L. J. Schurgers, Y. T. van der Schouw and J. W.
690 J. Beulens, The effect of menaquinone-7 supplementation on vascular calcification in
691 patients with diabetes: a randomized, double-blind, placebo-controlled trial, *Am. J.*
692 *Clin. Nutr.*, 2019, **110**, 883-890.
- 693 82. V. M. Brandenburg, S. Reinartz, N. Kaesler, T. Krüger, T. Dirrichs, R. Kramann, F.
694 Peeters, J. Floege, A. Keszeci and N. Marx, Slower progress of aortic valve
695 calcification with vitamin K supplementation: results from a prospective
696 interventional proof-of-concept study, *Circulation*, 2017, **135**, 2081-2083.
- 697 83. M. K. Shea, C. J. O'Donnell, U. Hoffmann, G. E. Dallal, B. Dawson-Hughes, J. M.
698 Ordovas, P. A. Price, M. K. Williamson and S. L. Booth, Vitamin K supplementation
699 and progression of coronary artery calcium in older men and women, *Am. J. Clin.*
700 *Nutr.*, 2009, **89**, 1799-1807.
- 701 84. L. A. J. L. M. Braam, A. P. G. Hoeks, F. Brouns, K. Hamulyák, M. J. W.
702 Gerichhausen and C. Vermeer, Beneficial effects of vitamins D and K on the elastic
703 properties of the vessel wall in postmenopausal women: a follow-up study, *Thromb.*
704 *Haemost.*, 2004, **91**, 373-380.
- 705

706 **Table 1:** A compilation of commonly consumed foods that contain vitamin K1 and/or K2 obtained from a range of existing vitamin K resources.

		Vitamin K content (µg/100 g or µg/100 ml)											
Type	Food	K1	MK4	MK5	MK6	MK7	MK8	MK9	MK10	MK11	MK12	MK13	Country ¹
Milk	Full cream milk	0.5	0.8	0.1	UD	UD	UD	UD	UD	-	-	-	NL ³
	Reduced fat milk	0.2	UD	-	-	-	-	-	-	-	-	-	USA ¹³
	Soya milk	3.0	-	-	-	-	-	-	-	-	-	-	USA ¹³
Fats & Oils	Margarine	93.2	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Butter	14.9	15	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Olive	53.7	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Butter & margarine blend	86.5	1.7	-	-	-	-	-	-	-	-	-	USA ¹³
	Canola	71.3	UD	-	-	-	-	-	-	-	-	-	USA ¹³
Cheese	Cheddar	2.4	9.5	UD	0.9	0.8	5.6	175	42.9	42.2	1.3	UD	USA ¹⁷
	Low fat cheddar	0.5	1.8	UD	UD	0.7	4.0	22.6	1.5	16.3	UD	UD	USA ¹⁷
	Cottage cheese	0.3	0.3	0.5	0.5	0.6	2.5	8	0.4	39.1	UD	UD	USA ¹⁷
	Brie	4.9	12.5	UD	UD	UD	UD	UD	UD	-	-	-	FRN ⁴⁰
	Camembert	2.5	8.0	1.3	0.1	3.2	1.5	4	UD	-	-	-	FRN ⁴⁰
	Parmesan	20.6	UD	UD	0.01	0.1	0.2	UD	UD	-	-	-	ITL ⁴⁰
	Cream cheese	2.4	UD	-	-	-	-	-	-	-	-	-	USA ¹³
Creams	Yogurt (regular full-fat)	0.4	0.7	UD	UD	UD	UD	13.2	1.6	8.4	UD	UD	USA ¹⁷
	Standard full cream	2.4	9.3	UD	UD	UD	UD	442	85.2	44.3	2.6	UD	USA ¹⁷
	Ice cream	0.3	UD	-	-	-	-	-	-	-	-	-	USA ¹³
Breads	White bread	3.4	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Multigrain bread	1.5	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Rye bread	0.7	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
Meats	Beef	0.02	1.4	UD	UD	1.3	3.7	UD	UD	-	-	-	NL ⁴⁰
	Minced Meat	1.1	7.6	UD	UD	UD	UD	UD	UD	-	-	-	NL ⁴⁰
	Deer	2.4	0.9	UD	UD	UD	UD	UD	UD	-	-	-	NL ⁴⁰
	Chicken	UD	10.1	UD	UD	UD	UD	UD	UD	-	-	-	NL ⁴⁰

	Pork	UD	1.4	UD	UD	UD	UD	UD	UD	-	-	-	NL ⁴⁰
	Salmon	1.3	5.7	UD	UD	UD	UD	UD	UD	-	-	-	NL ⁴⁰
	Salami	2.3	9	UD	UD	UD	UD	UD	UD	-	-	-	NL ⁴⁰
	Goose liver	10.9	369	UD	UD	UD	UD	UD	-	-	-	-	NL ⁴⁰
	Egg yolk	2.1	31.4	UD	0.7	UD	UD	UD	UD	-	-	-	NL ⁴⁰
	Bacon	0.3	3	UD	UD	UD	UD	UD	2.0	29.7	UD	UD	USA ¹⁷
	Lamb	4.5	UD	-	-	-	-	-	-	-	-	-	USA ¹³
	Ham	UD	4.3	-	-	-	-	-	-	-	-	-	USA ¹³
	Pork sausage	0.3	18.3	-	-	-	-	-	-	-	-	-	USA ¹³
	Tinned tuna (in water)	2.5	0	-	-	-	-	-	-	-	-	-	USA ¹³
Fruits	Apples	3.0	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Pears	5.2	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Avocados	21	-	-	-	-	-	-	-	-	-	-	USA ¹³
Vegetables	Kale	817.0	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Broccoli	156.0	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Spinach	387.0	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Peas	36.0	UD	UD	UD	UD	UD	UD	UD	-	-	-	NL ³
	Cucumber	16.4	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Lettuce (green)	126.3	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Lettuce (red)	140.3	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Celery	29.3	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Carrots	13.2	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Cabbage	76.0	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Cauliflower	20.2	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Potatoes	2.0	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Tomatoes	7.9	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Capsicum (green)	7.4	-	-	-	-	-	-	-	-	-	-	USA ¹³
Zucchini	4.2	-	-	-	-	-	-	-	-	-	-	USA ¹³	
Kidney beans	19.0	-	-	-	-	-	-	-	-	-	-	USA ¹³	

	Cashews	34.1	-	-	-	-	-	-	-	-	-	-	USA ¹³
	Tofu (firm)	12.0	0.01	-	-	UD	-	-	-	-	-	-	JPN ¹⁵
Fermented vegetables	Natto	32.1	UD	7.2	12.4	996.5	82.4	UD	UD	-	-	-	NL ⁴⁰
	Sauerkraut	22.4	0.4	0.9	1.6	0.2	0.9	1.5	UD	-	-	-	NL ⁴⁰

707 ¹ indicates origin of produce and/or country where vitamin K measurement was performed followed by reference. UD: undetected; -: unknown; USA: United States of
708 America; NL: the Netherlands, JPN: Japan; FRN: France; ITY: Italy

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