The role of menaquinones (vitamin K2) in human health

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Abstract

Recent reports have attributed the potential health benefits of vitamin K beyond its function to activate hepatic coagulation factors. Moreover, several studies have suggested that menaquinones, also known as vitamin K₂, may be more effective in activating extra-hepatic vitamin K-dependent proteins than phylloquinone, also known as vitamin K₁. Nevertheless, present dietary reference values (DRV) for vitamin K are exclusively based on phylloquinone, and its function in coagulation. The present review describes the current knowledge on menaquinones based on the following criteria for setting DRV: optimal dietary intake; nutrient amount required to prevent deficiency, maintain optimal body stores and/or prevent chronic disease; factors influencing requirements such as absorption, metabolism, age and sex. Dietary intake of menaquinones accounts for up to 25% of total vitamin K intake and contributes to the biological functions of vitamin K. However, menaquinones are different from phylloquinone with respect to their chemical structure and pharmacokinetics, which affects bioavailability, metabolism and perhaps impact on health outcomes. There are significant gaps in the current knowledge on menaquinones based on the criteria for setting DRV. Therefore, we conclude that further investigations are needed to establish how differences among the vitamin K forms may influence tissue specificities and their role in human health. However, there is merit for considering both menaquinones and phylloquinone when developing future recommendations for vitamin K intake.

Key words: Vitamin K; Dietary recommendations; Menaquinones; Bone health; CVD; Bioavailability

There is increasing interest in the potential health benefits of vitamin K beyond its role in coagulation. Several studies have reported functions for vitamin K beyond its classic role, including the improvement of bone health¹¹, and the reduction of vascular calcification and cardiovascular risk²⁻³. Moreover, several studies²⁻⁴ have suggested that menaquinones, also known as vitamin K₂, could be more effective in these functions than phylloquinone, also known as vitamin K₁. Nevertheless, menaquinones are generally not taken into consideration when developing dietary recommendations for vitamin K. Present recommendations for dietary vitamin K are defined for phylloquinone intake only, and are based on median intakes of phylloquinone in certain regions, such as North America⁵⁻⁷. In some cases, the effects of phylloquinone on coagulation have also been accounted for. For healthy adults, adequate intakes of vitamin K range from 55 to 90 μg/d for adult women and 65–120 μg/d for adult men.

The International Life Sciences Institute (ILSI) Europe has selected experts on vitamin K from academia and industry to review the need for specific dietary reference values (DRV) for menaquinones. To achieve this objective, the expert group conducted a thorough review of existing literature on

Abbreviations: BMD, bone mineral density; DRV, dietary reference values; Gla, γ-carboxyglutamate; ILSI, International Life Sciences Institute; MGP, matrix γ-carboxyglutamate protein; MK-n, menaquinones; OC, osteocalcin; PIVKA-II, undercarboxylated prothrombin; PT, prothrombin time; ucOC, uncarboxylated osteocalcin.

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dietary menaquinones and their role in human health to evaluate: (a) whether unique recommendations for menaquinone intake are justified at this time and (b) what additional information is needed to have strong scientific underpinnings for establishing DRV for menaquinones. In defining nutrient requirements, the selection of criteria to establish nutrient adequacy is an important step. For most nutrients, a hierarchy of criteria for nutrient adequacy can be established, ranging from the prevention of clinical deficiency to the maximisation of body stores or status. The goal is to have a low probability of nutrient inadequacy while minimising the potential risk of excess. In light of this definition, we have reviewed the literature with a focus on the following criteria for setting DRV: chemical structure and function of menaquinones; dietary intake of menaquinones; absorption and metabolism of menaquinones; amount of menaquinones required to prevent deficiency; maintain optimal body stores and/or prevent chronic disease; factors influencing menaquinone requirements such as age, sex and safety. The evidence for individual menaquinones for each of these items is described and, if known, differences with phylloquinone are described. Based on this evidence, a conclusion on setting a DRV for menaquinones is drawn and recommendations for future research are made.

Chemical structure and function of menaquinones

Vitamin K is a generic term for a number of structurally related compounds that are characterised by their common functional methylated naphthoquinone ring system, and an aliphatic side chain composed of a number of isoprenoid residues. All differences between the various forms of vitamin K originate from the differences in the length and the saturation degree of the side chain. Phylloquinone is a single compound with a side chain of four isoprenoid residues, three of which are saturated (Fig. 1). Menaquinones, commonly found in nature, have side chains of varying length between four and thirteen isoprene residues, most of which are unsaturated. However, some bacteria produce isoprenologues in which one or more of the prenyl units are saturated. Menaquinones are generally denoted as MK-\(n\), where \(n\) stands for the number of isoprene residues.

MK-4 is unique among the menaquinones in that it is not synthesised by bacteria. Instead, MK-4 is alkylated from menadione (vitamin K\(_3\), a synthetic form of vitamin K that is present in animal feeds, or is the product of tissue-specific conversion directly from dietary phylloquinone, with menadione as the postulated intermediate. There is also speculation that longer-chain menaquinones, such as MK-7, can be converted to MK-4 as well. The most abundant menaquinones in the human diet are the short-chain MK-4, which originates from animal products, and the long-chain MK-7, MK-8, MK-9 and MK-10.

All forms of vitamin K have one well-known function. They all serve as a cofactor for the post-translational enzyme \(\gamma\)-glutamyl carboxylase, which is established by the common naphthoquinone ring structure. This enzyme converts certain protein-bound glutamate residues into \(\gamma\)-carboxyglutamate, generally known as Gla. Currently, seventeen members of the Gla protein family are known, including seven proteins involved in blood coagulation (all synthesised in the liver), osteocalcin (OC; bone), matrix Gla protein (MGP; mainly cartilage and vessel wall), growth arrest-specific protein 6, Gla-rich proteins, two proline-rich Gla proteins, two transmembrane Gla proteins, peristin and peristin-like factor. With the exception of the clotting factors OC (bone formation) and MGP (inhibitor of soft tissue calcification), the physiological importance of these proteins is not yet fully understood. At this time, conformation-specific assays are available for two extra-hepatic Gla proteins (OC and MGP).

Dietary intake of menaquinones

Menaquinones generally are of microbial origin. Important dietary sources are cheese, cured and natto (a traditional Japanese food composed of fermented soya beans), while dietary phylloquinone is mainly found in green vegetables, notably spinach, broccoli, kale and Brussels sprouts. Estimated intake of phylloquinone and menaquinones in The Netherlands and Germany has suggested that between 10 and 25% of total vitamin K intake are provided by menaquinones from cheese, curd and natto, while long-chain menaquinones, MK-7 to MK-10, are predominantly consumed by high dairy intake populations, such as the Dutch population. In addition, cheese is the most important source of menaquinones in the European food supply, and therefore menaquinone concentrations in the cheeses of Dutch, German, Swiss, British and French origins were tested. Since different lactic acid bacteria are used in European cheeses, a large variability in

![Fig. 1. Chemical structures of K vitamins. MK, menaquinone.](https://www.cambridge.org/core/journals/british-journal-of-nutrition/article/chemical-structure-and-function-of-menaquinones/583F22B24C4D1F680B15F3D470C0210F)
menaquinone content among the cheeses was found. However, it remains to be seen whether these data are applicable to non-European countries.

The few studies that have provided estimates for menaquinone intakes have been mainly performed among Japanese or European populations for which menaquinone-rich foods are present in the diet. A study by Kamao et al. measured the menaquinone content of foods and the estimated intake of MK-4 and MK-7 of 125 young Japanese women using a 3 d weighed food record. The MK-7 intake of this population was originally estimated at 57.4 (SD 83.7) μg/d and accounted for approximately 25% of the total intake of vitamin K. However, this estimate is mainly driven by natto consumption, as it accounted for 99% of the MK-7 intake and almost half of the population studied consumed natto. After stratifying by natto consumption, intake of MK-7 was estimated at 154.1 (SD 87.8) μg/d among natto consumers, but data on non-natto consumers were not provided.

Several Dutch studies investigating the associations of menaquinone intake with disease incidence obtained estimates for menaquinone intake(2,3,22,23) (Table 1) using FFQ. These intake estimates were based on direct measurements of menaquinones in foods in combination with published data(16,24). The self-reported mean intake of MK-7 of this population was estimated at 31.6 (SD 12.3) μg/d and accounted for approximately 25% of the total intake of vitamin K. In the EPIC-Netherlands cohort, cheese contributed 53% of menaquinone intake, while milk products and meat contributed 19 and 17%, respectively. The most prominent long-chain menaquinone reported in the diet was MK-9. In the Rotterdam Study, MK-5 to MK-10 contributed 23.1 (SD 16.3) μg/d for men and 20.7 μg/d for women. These data on food contents of menaquinones have also been applied to a German cohort of approximately 12,000 men(19). Similar estimates of 34.7 μg/d (interquartile range 25.7–45.7) were reported for all menaquinones and 18.0 μg/d (11.7–27.0) for MK-5 to MK-9, with cheese being the most important food source of menaquinones(19).

Finally, the European Food Safety Authority reported data on UK intake of menaquinones based on the UK National Dietary and Nutrition Survey(25). This study used weighed dietary records, albeit based on the menaquinone content from a limited number of food items(25). The overall estimated intake of menaquinones ranged from 36 (female adults) to 54 (male teenagers) μg/d. The mean estimated intake of menaquinones among male adults was 43 μg/d, which is similar to the intakes reported for other European countries.

It should be noted that estimates from the Dutch and German populations were obtained from FFQ that are designed to estimate the relative dietary intake of large populations, but not to estimate the absolute dietary intake. These limitations should be kept in mind when interpreting these data. In order to obtain more precise estimates of menaquinone intakes, studies using (weighed) food records and concentrations of individual menaquinones obtained from representative foods from different food supplies are required. Nonetheless, current studies have shown estimated intakes

<table>
<thead>
<tr>
<th>Study</th>
<th>Participants</th>
<th>Location</th>
<th>Dietary assessment</th>
<th>Menquinone intake (μg/d)</th>
<th>Endpoint</th>
<th>BAC, breast arterial calcification; CAC, coronary artery calcification.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maas et al.</td>
<td>1689 women, aged 49–70 years</td>
<td>The Netherlands</td>
<td>FFQ</td>
<td>26·9 BAC → 29·4 BAC</td>
<td>Coronary calcification</td>
<td>90 (0.65–0.96)</td>
</tr>
<tr>
<td>Beulens et al.</td>
<td>564 postmenopausal women, aged 49–70 years</td>
<td>The Netherlands</td>
<td>FFQ</td>
<td>Mean: 31·6 (SD 12·3) μg/d</td>
<td>CAC</td>
<td>0·80 (0·65–0·98)</td>
</tr>
<tr>
<td>Geleijnse et al.</td>
<td>4473 men and women, aged 55 years and over</td>
<td>The Netherlands</td>
<td>FFQ</td>
<td>Men: 30·8 (SD 18·0) μg/d; women: 27·0 (SD 15·1) μg/d</td>
<td>CAC</td>
<td>0·48 (0·32–0·71)</td>
</tr>
<tr>
<td>Gast et al.</td>
<td>4977 adults, aged 55 years and over</td>
<td>The Netherlands</td>
<td>FFQ</td>
<td>Mean: 29·1 (SD 12·8) μg/d</td>
<td>CHD</td>
<td>0·91 (0·65–1·00) per 10 μg increment</td>
</tr>
</tbody>
</table>

Table 1. Menaquinone intake, arterial calcification and risk of CHD
of menaquinones ranging between 30 and 50 µg/d, which account for up to 25% of intake of total vitamin K.

Absorption and metabolism of menaquinones

Phylloquinone is primarily obtained from green, leafy vegetables in which it is tightly bound to the membranes of plant chloroplasts, and thus less bioavailable compared with phylloquinone obtained from plant oils and/or dietary supplements(13). Menaquinones, which are primarily derived from animal-based sources, are consumed in food matrices containing more fat that may improve absorption and lead to higher bioavailability than phylloquinone(26). However, this has yet to be systematically tested for all menaquinones.

Following intestinal absorption, all vitamin K forms are incorporated into TAG-rich lipoproteins and transported primarily to the liver, but also to other target tissues. Circulating TAG-bound forms of vitamin K peak at around 4–10 h after intake and the majority of phylloquinone and MK-4 are removed from the circulation by 24 h postprandially(13,16,27).

Currently, human data on the absorption of menaquinones from food sources are limited to MK-7. These MK-7 data show similar peaks at 4 h after intake, but MK-7 does not appear to be completely removed from the circulation after 72–96 h(15,16). The different pharmacokinetics among various vitamin K forms also result in very different plasma half-life times. Whereas phylloquinone has a relatively short half-life time(26), MK-7 has a reported half-life time of several days(15,27).

Available data indicate higher absorption and bioavailability of MK-7 than phylloquinone, which may facilitate its uptake by various target tissues. Another difference between the short-chain forms of vitamin K (phylloquinone and MK-4) and the long-chain forms relates to tissue distribution. For example, one study showed that MK-9 is preferentially incorporated in LDL, which facilitates its transport to non-hepatic target tissues(29).

It is not known whether other long-chain menaquinones have similar transport differences. MK-4 is unique among the menaquinones in its tissue distribution, which relates to its non-bacterial origin. As recently shown in a rodent model using MK-4, the labelled MK-4 was most abundant in the brain, kidney, fat and reproductive organs. In contrast to phylloquinone as the sole dietary source of vitamin K, there was no conversion of phylloquinone to MK-4 in the liver nor were there detectable amounts of labelled MK-4 in serum. These data confirm earlier rodent studies that have reported differences in tissue distribution between phylloquinone and MK-4(30,31).

A caveat to these conclusions is that the data for phylloquinone are much more comprehensive than those for menaquinones. Many investigators have studied phylloquinone pharmacokinetics using different study designs, including stable isotopes(32–34). In contrast, menaquinone pharmacokinetic data are limited: (1) in the forms studied (mainly limited to MK-7); (2) from a lack of replication by independent laboratories; (3) by an absence of using stable isotope technology. More research is clearly required to quantify the differences in absorption and bioavailability among the various forms of vitamin K in order to set nutrient requirements.

Microbiotic production of menaquinones

Most aerobic Gram-positive bacteria and the majority of anaerobic bacteria produced by the gut use menaquinones in their electron transport pathways. The length of the side chain, as indicated by different menaquinones, is controlled by specific bacteria(10). The reasons for this are not entirely clear, but the length and degree of saturation of the menaquinone side chain are often dependent on the growth temperature of a given species(50). Based on qualitative bacteriological analyses, several bacteria have been identified to produce specific menaquinones. Menaquinones produced by the gut flora have been tabulated by previous studies(37–39). For example, Bacteroides fragilis produces MK-10 to MK-12, while Eubacterium lentum produces MK-6. Likewise, bacteria used as starter cultures for the production of foods such as cheese may also produce specific menaquinones. For example, Lactococcus lactis ssp. lactis and L. lactis ssp. cremoris produce mainly MK-8 and MK-9(40), while propionibacteria produce mainly MK-9(41).

The implications for the relative bioavailability of dietary menaquinones produced by bacteria in the food supply need to be considered relative to the bioavailability of menaquinones produced by bacteria in the human intestine.

It was once stated that up to 50% of the human requirement for vitamin K was fulfilled by the intestinal production of menaquinones(42,43). The 50% estimate was based on semi-quantitative measurements of the vitamin K content of the human liver, in which one-half of the vitamin K content was phylloquinone and the other half was a mixture of long-chain menaquinones(43). However, subsequent studies indicated that phylloquinone accounted for less than 10% of the vitamin K content in the human liver, with a greater preponderance of MK-10, MK-11 and MK-12 than previously assumed(43,45).

Based on these hepatic menaquinone concentrations, one would predict that circulating long-chain menaquinones would be in much higher concentrations than phylloquinone should these menaquinones have a major contribution to the human requirement for vitamin K. This, however, does not appear to be the case, and the route of absorption of bacterially produced menaquinones is still unclear. The absorption of all vitamin K forms takes place in the small intestine via a process requiring bile salts(46). Bile salts are absent in the colon where the majority of menaquinones are produced, suggesting a low absorption of these vitamin K forms(47). This was confirmed by Ichihashi et al.(48), who showed that the absorption of intestinally produced menaquinones in rats is low and that the absorption rates decrease markedly with the length of the side chain. A study in infants also indicated that intestinally produced menaquinones are not well absorbed(49). This study compared faecal and serum concentrations of phylloquinone and menaquinones of formula-fed infants with breast-fed infants. Formula-fed infants had higher serum and faecal phylloquinone concentrations as well as a higher MK-5 to MK-9 faecal concentration.
compared with breast-fed infants. Serum menaquinones were undetected in most formula-fed infants, suggestive of poor absorption\(^1\). Another consideration is that most bacterially produced menaquinones are within the bacterial membranes, hence not readily bioavailable. It has been postulated that these bacterially synthesised menaquinones may be important in maintaining normal coagulation among severely ill patients with prolonged vitamin K deficiency\(^2\); however, current data are inconclusive regarding the relative contribution of menaquinones to fulfilling the dietary requirements for vitamin K.

### The amount of menaquinones required to prevent deficiency and maintain optimal body stores

To understand the impact of menaquinones on health, it is necessary to demonstrate the link between the intakes of menaquinones and the nutritional status of vitamin K. Several biochemical markers of vitamin K status are available and all have their strengths and weaknesses, as detailed elsewhere\(^3\). However, measures of plasma or tissue menaquinone concentrations are needed to isolate the effects of menaquinones from those of phylloquinone on human health. Other markers of vitamin K status, which include urinary metabolites of vitamin K\(^4\), coagulation times and uncarboxylated Gla proteins, cannot differentiate the effects of menaquinones from phylloquinone. Therefore, differences between menaquinones and phylloquinone can only be determined through the use of study designs that directly compare the response of individual biomarkers with the intakes of individual forms of vitamin K.

Under controlled conditions of dietary intake, circulating menaquinone concentrations increase in response to the high intake of menaquinones and decline over time when the dietary source of menaquinones is removed\(^5\). However, data are limited, since the HPLC and MS techniques are limited to a few qualified laboratories and the long-chain menaquinones are often below the detection limit in the circulation when measured in the general population. Only a few studies have measured plasma menaquinones in response to the intake of individual menaquinones, and these have been limited to MK-7 or MK-4 supplementation\(^6\). For plasma MK-7, two studies showed a clear dose–response effect on circulating MK-7 concentrations after supplementation with doses ranging between 45 and 420 \(\mu g\) \(^7\). In contrast, MK-4 was not detected in the circulation following a single dose of 420 \(\mu g\) \(^8\).

Only two studies have investigated the response of vitamin K urinary metabolites to single oral doses of menaquinones. Both studies showed a good response of urinary menadione\(^9\) or side-chain catabolite\(^10\) excretion to relatively high doses of 15 or 45 mg of MK-4 or 1 mg of MK-7. Both studies included a direct comparison of menaquinones with phylloquinone, and showed similar results for both vitamin K forms. Harrington et al.\(^11\) showed that the excretion of the 5- and 7-carbon side-chain metabolites responds to the depletion and repletion of phylloquinone. A similar response to menaquinones would be expected, but no studies of similar design have been conducted.

Prothrombin time (PT), also expressed as an international normalised ratio, is a test of coagulation that can reflect clinical deficiency of vitamin K due to frank deficiency or the antagonism of vitamin K. However, PT is non-specific because abnormal values are also indicative of diseases unrelated to vitamin K deficiency. PT changes only when prothrombin concentrations drop below 50% of normal, demonstrating its low sensitivity for detecting the deficiency of vitamin K\(^12\). To date, only two studies\(^13,15\) have reported the effects of MK-7 and MK-9 on coagulation parameters. In these studies, the antidotal effect of single doses of MK-7 and MK-9 was studied in volunteers stabilised on oral vitamin K antagonists. Both studies showed that MK-7 and MK-9 decreased the international normalised ratio and the concentrations of coagulation factors, and this effect was stronger for MK-7 than phylloquinone\(^13,15\). However, in one study\(^7\), menaquinones were provided as different food sources with differing doses and bioavailability, which may have influenced the results. Although these studies are informative in the clinical context of the reversal of oral anticoagulation, they are not suitable for determining the amount of menaquinones required to prevent deficiency or maintain optimal body stores. Although the effects of coagulation factors on depletion or repletion with menaquinones have not been investigated to date, sustained intakes as low as 10 \(\mu g\)/d of phylloquinone for several weeks do not prolong PT in otherwise healthy adults\(^7,59–60\).

Thus far, conformation-specific tests have been developed for prothrombin, OC and MGP to evaluate the extent to which the various Gla proteins are carboxylated in healthy subjects. Advantages of measuring uncarboxylated vitamin K-dependent proteins are that insufficiencies measured in circulating forms theoretically reflect what occurs at the tissue level. Undercarboxylated prothrombin, also known as PIVKA-II, detects abnormalities in prothrombin before the prolongation of PT, but does not have the sensitivity to detect the variability of usual vitamin K intake observed in normal healthy populations. PIVKA-II has been used as a marker of vitamin K status in healthy people and has been shown to respond to both dietary depletion and subsequent repletion with phylloquinone\(^61,62\). However, only one study\(^63\) investigated the effect of a single intravenous dose of 10 mg MK-4 in vitamin K-deficient cancer patients, and showed a decrease in PIVKA-II levels 1–3 d after ingestion.

The effect of menaquinones on the proportion of OC that is uncarboxylated (ucOC) has been more frequently studied. ucOC is highly responsive to supplementation with either MK-4 or MK-7 in doses ranging from 45 \(\mu g\)/d to 45 mg/d (MK-4 only)\(^13,54,64–66\). Only a low dose of 45 \(\mu g\)/d MK-7 did not lead to a significant reduction in ucOC\(^54\). A direct comparison of phylloquinone with MK-7 supplementation indicated that MK-7 is more effective in carboxylating OC than phylloquinone\(^15\). Assays to measure desphospho-uncarboxylated MGP only recently became available. Since that time, several intervention studies have shown clear dose–response effects of desphospho-uncarboxylated MGP...
to MK-7 supplementation with doses ranging between 10 and 360 μg/d\(^{67–70}\). However, a direct comparison between menaquinones and phylloquinone in the response of desphospho-uncarboxylated MGP to supplementation by the individual vitamin K forms has not been made.

The amount of menaquinones required to prevent chronic diseases

**Menaquinones, coronary calcification and CVD**

Coronary artery calcification is an important predictor of CVD\(^{71}\). MGP is an inhibitor of vascular calcification\(^{72}\). Through carboxylation of MGP, vitamin K may help reduce coronary calcification and thereby reduce the risk of CVD. Observational studies have indeed shown that a high intake of vitamin K is associated with reduced coronary calcification and a reduced risk of CVD\(^{2,3,23}\). The results from some studies suggest that this is mainly due to menaquinones\(^{2–4,23}\). Thus far, three cross-sectional studies\(^{2,5,73}\) investigated the associations of menaquinone intake and coronary calcification, as summarised in Table 1. In the Rotterdam Study, intakes of menaquinones were lower in participants with severe aortic calcifications (25.6 μg/d) than in participants with moderate or mild calcifications (28.6 and 28.8 μg/d, respectively; \(P=0.001\))\(^{73}\). A strong inverse relationship between menaquinone intake and severe calcification was found in the mid and upper tertiles of menaquinone intake compared with the lowest tertile, reaching significance in the highest tertile with a menaquinone intake of more than 32.7 μg/d. Using breast arterial calcification as a measure of arterial calcification, the prevalence of breast arterial calcification was less common in the highest (9%) quartile of menaquinone intakes, compared with the lowest quartile (13%)\(^{75}\). This study showed a similar association to that of Geleijnse et al.\(^{53}\) with an OR of 0.7 (95% CI 0.5, 1.1), although it did not reach significance. Similarly, a high menaquinone intake over 48 μg/d was associated with reduced coronary calcification among 600 middle-aged women\(^{76}\). We are not aware of any randomised trials to date that investigated the effect of menaquinones on the progression of arterial calcification.

Also, two of the previously mentioned cohort studies investigated the relationship between menaquinone intake and the risk of CHD (Table 1). In the Rotterdam cohort\(^{53}\), the relative risk of incident CHD was reduced in the upper tertile of menaquinone intake compared with the lowest tertile (0.43; 95% CI 0.24, 0.77). In the Prospect-EPIC cohort\(^{2,3}\), the investigators also observed an inverse association between the intake of menaquinones and the risk of CHD with a hazard ratio of 0.51 (95% CI 0.50, 1.08) per 10 μg/d of menaquinone intake. In order to compare these results with previous studies using categories, a menaquinone intake of 35 μg/d would lead to a hazard ratio of about 0.7, which compares nicely with previous studies. The association between menaquinone intake and the incidence of stroke has not been investigated to date. Of note, several of these studies also investigated the relationship between phylloquinone intake and coronary calcification or the risk of CHD, but could not detect significant associations\(^{2,5,23}\). Whether this is due to biological differences between menaquinones and phylloquinone or perhaps lower validity of the FFQ to estimate phylloquinone intake is currently unclear\(^{23}\). Finally, these associations have only been investigated in Dutch populations and generalisability of these results should be studied in different populations.

**Menaquinones and bone**

In bone, three vitamin K-dependent proteins have been isolated: protein S; MGP; OC. The anticoagulant protein S is synthesised by osteoblasts (bone-forming cells), but its role in bone metabolism is unclear. MGP has been found in bone, dentine, cartilage and soft tissue, including blood vessels, and is associated with the organic matrix and mobilisation of bone Ca. The results of animal studies suggest that MGP prevents the calcification of soft tissue and cartilage, while facilitating normal bone growth and development\(^{24}\). OC is a protein synthesised by osteoblasts. The synthesis of both OC and MGP is regulated by calcitriol and retinoic acid. Higher ucOC concentrations, indicating a low vitamin K status, were associated with a higher hip fracture risk and lower bone mineral density (BMD) in adults\(^{75–85}\) and children\(^{86–89}\). Unfortunately, the proportion of ucOC does not differ between phylloquinone and menaquinones in terms of the form responsible as an enzyme cofactor. However, serum menaquinone levels were lower in patients with osteoporosis, osteopenia and osteoporotic fractures compared with controls\(^{90–93}\). In addition, an inverse association was found between circulating MK-7 levels and the incidence of vertebral fractures in Japanese women, although this association was stronger for phylloquinone and fracture risk\(^{94}\).

Intervention studies using pharmacological doses of MK-4 showed beneficial effects on bone parameters\(^{1}\). However, these intervention studies used very high doses of MK-4 (generally 45 mg) that cannot be obtained from the habitual diet. Since this paper is focused on dietary doses that are relevant for nutritional requirements, we will focus on intervention studies that used doses that can be nutritionally obtained. Although natto contains more than 100 times as much menaquinones as various cheeses, studies on natto or its equivalent amount of MK-7 are considered within the dietary intake range.

Intervention studies investigating the effect of menaquinone supplementation on bone markers are shown in Table 2. Whereas MK-7 at low doses did not affect bone formation\(^{64,65}\), intake of natto three times per week increased bone-specific alkaline phosphatase when compared with once-per-week natto intake\(^{95}\). Only one study showed decreased bone resorption due to a combination of vitamin K forms, vitamin D\(_3\), Ca and lifestyle recommendations\(^{96}\). This finding was independent of the form of vitamin K taken, although on a molecular basis, the daily intake of MK-7 (0.154 μmol) in this study was about 30% less than that of phylloquinone (0.221 μmol).

A few cross-sectional studies investigated the association between menaquinone intake and bone maintenance. For
Table 2. Intervention studies on the dietary levels of vitamin K, bone markers and bone mineral density (BMD)

<table>
<thead>
<tr>
<th>Study design</th>
<th>Participants</th>
<th>Location</th>
<th>Duration (months)</th>
<th>Intervention</th>
<th>BMD</th>
<th>Bone markers</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>DBPC trial</td>
<td>173 women aged 55–65 years</td>
<td>Greece</td>
<td>12</td>
<td>Four groups: (1) control group; (2) dairy product with Ca and vitamin D; (3) phylloquinone; (4) dairy product with MK-7 for 12 months</td>
<td>No difference</td>
<td>No difference</td>
<td>Unaffected†‡</td>
</tr>
<tr>
<td>DBPC trial</td>
<td>Fifty-five Dutch children aged 6–10 years</td>
<td>The Netherlands</td>
<td>12</td>
<td>Four groups: (1) placebo; (2) placebo and MK-7; (3) 360 mg MK-7 for 12 weeks; (4) three times per week natto intake at lunch</td>
<td>No difference</td>
<td>No difference</td>
<td>Unaffected†‡</td>
</tr>
<tr>
<td>DBPC trial</td>
<td>Seventy-three postmenopausal women aged 50–60 years</td>
<td>Japan</td>
<td>12</td>
<td>Four groups: (1) no natto intake; (2) once per month; (3) once per week; (4) three times per week natto intake at lunch</td>
<td>No difference</td>
<td>Significant higher concentration group (3)</td>
<td>Significant higher concentration group (3)†‡</td>
</tr>
</tbody>
</table>

Bone markers: Decreased in groups (3) and (4) compared with the other groups*.

DBPC, double-blind placebo-controlled; MK-7, menaquinone-7; NA, not applicable.

* Urinary deoxypyridinoline.
† Bone-specific alkaline phosphatase.
‡ Degradation products of C- or N-terminal telopeptides of type I collagen.

Example, two Japanese studies showed that the usual dietary intake of natto was effective in maintaining bone stiffness(97) and was positively associated with a 3-year change in BMD at the femoral neck(98). Within Norwegian individuals, no linear association was found between dietary menaquinones and BMD of the total hip, but there were even less data for menaquinones. Currently, data on the intake or supplementation of menaquinones are limited to the assessment in children and teenagers in the UK National Dietary and Nutrition Survey(25).

Requirements across the life cycle

Dietary intake has historically been considered the primary determinant of vitamin K status(11). However, other non-dietary factors are emerging as determinants, such as age and ethnicity(51). To develop recommendations for dietary intakes(9), sufficient data are needed to evaluate requirements across the life cycle. The data for phylloquinone are sparse(7), but there are even less data for menaquinones. Currently, data on the intake or supplementation of menaquinones are limited to the assessment in children and teenagers in the UK National Dietary and Nutrition Survey(25).

The only clinically indicated use of vitamin K is as a prophylactic against vitamin K deficiency bleeding in otherwise healthy-appearing neonates(102). The low content of vitamin K in breast milk, low placental transfer of vitamin K, low levels of clotting factors at birth and a sterile gut are all contributing factors to the risk of vitamin K deficiency bleeding in the first few months of life. Prevention of vitamin K deficiency bleeding by oral or intramuscular administration of vitamin K at birth is standard practice in many countries. Whereas most countries use phylloquinone, certain Asian countries,
including Japan, use MK-4 prophylactically\textsuperscript{102}. At no other point in the life cycle is frank deficiency of vitamin K a concern among an otherwise healthy population.

Safety of high vitamin K intake

There is no documented case of toxicity for phylloquinone or menaquinones\textsuperscript{7,25}. The European Food Safety Authority’s safety assessment of menaquinones as a source of vitamin K added for nutritional purposes concluded that low doses of menaquinones presented no safety concerns\textsuperscript{25}. Similarly, an animal study reported no toxicity associated with synthetic MK-7 administered in a single oral dose up to 2000 mg/kg or for 90 d of oral administration of 10 mg/kg per d\textsuperscript{103}.

It is often postulated that excessive vitamin K may result in overcoagulation, i.e. increased thrombosis risk. However, vitamin K-dependent proteins have a limited number of Glu residues capable of γ-carboxylation per molecule, beyond which there can be no further γ-carboxylation or excessive coagulation. Despite this, it is critical to demonstrate that a high intake of menaquinones does not increase thrombosis risk. It was shown in rats that thrombosis risk is not increased at doses up to 250 mg/kg of MK-4\textsuperscript{104}. In human subjects, the endogenous thrombin potential, which is the most sensitive marker to evaluate thrombosis risk in plasma\textsuperscript{105}, was not increased at MK-4 intakes as high as 360 μg/d for 6 weeks\textsuperscript{700}. The only exception to this is observed in individuals on coumarin-based oral anticoagulants, for whom dietary supplementation with vitamin K can influence the stability of the international normalised ratio\textsuperscript{59,106}. MK-7 has the potential to interfere with oral anticoagulants at doses greater than 50 μg/d\textsuperscript{131}. However, there is little collective experience on the potential toxicity or adverse events associated with sustained menaquinone supplementation among individuals with normal coagulation.

Table 3. Summary and recommendations for future research

<table>
<thead>
<tr>
<th>Criterion</th>
<th>Difference with phylloquinone</th>
<th>Recommendation for future research</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structure</td>
<td>Length and degree of saturation of the side chain</td>
<td>Do long-chain menaquinones convert to MK-4?</td>
</tr>
<tr>
<td>Function</td>
<td>MK-4 has functions that may be unrelated to the role as an enzyme cofactor</td>
<td>Determine the location and biochemical pathways required for the conversion of phylloquinone to MK-4</td>
</tr>
<tr>
<td>Dietary intake</td>
<td>10–25 % of total intake of vitamin K</td>
<td>More accurate data on the food content of menaquinones over more countries</td>
</tr>
<tr>
<td>Absorption</td>
<td>Absorption of certain menaquinones higher than phylloquinone</td>
<td>Stable isotope studies to quantify relative differences in absorption among different menaquinones and with phylloquinone</td>
</tr>
<tr>
<td>Metabolism</td>
<td>Distribution to extra-hepatic tissue may differ with phylloquinone</td>
<td>Stable isotope studies to quantify relative differences with phylloquinone</td>
</tr>
<tr>
<td>Bioavailability</td>
<td>Longer half-life time of certain menaquinones</td>
<td>Stable isotope studies to quantify differences with phylloquinone</td>
</tr>
<tr>
<td>Effect on the status of vitamin K</td>
<td>None reported</td>
<td>More elaborate validation of biomarkers for the status of vitamin K</td>
</tr>
<tr>
<td>Effect on health outcomes</td>
<td>Stronger associations with coronary calcification and the risk of CHD</td>
<td>Studies with clinical endpoints to isolate the putative effects of individual menaquinones</td>
</tr>
<tr>
<td>Requirement across the life cycle</td>
<td>Both forms used prophylactically</td>
<td></td>
</tr>
<tr>
<td>Safety</td>
<td>None reported</td>
<td></td>
</tr>
</tbody>
</table>

MK-4, menaquinone-4.

In Asia, MK-4 is routinely used for osteoporosis treatment in doses of 45 mg/d without reported toxicity. Reported adverse effects associated with these high doses are limited to skin rashes that subside with cessation of the MK-4 dosing\textsuperscript{107}. As concluded by the European Food Safety Authority\textsuperscript{25}, phase I clinical trials have not yet been designed to test the safety of menaquinones nor has any form of vitamin K been adequately tested for mutagenicity. However, it is biologically implausible to attain such high levels of intake and sufficient bioavailability from menaquinones obtained from food sources to present a risk to health.

Conclusions

There is growing speculation that certain dietary menaquinones, while consumed in lower quantities than phylloquinone, may have unique and important contributions to the role of vitamin K on human health. However, present DRV for vitamin K are exclusively based on phylloquinone. In recognition of this emerging paradigm shift in vitamin K nutrition research, we have reviewed existing literature to evaluate the current state of knowledge on menaquinones that would be needed for inclusion in the DRV for vitamin K. It was concluded that differences in the chemical structure of menaquinones compared with phylloquinone may lead to differences in absorption and bioavailability (Table 3). Several studies have shown that certain forms of menaquinones may be more bioavailable and effective in carboxylating particular extra-hepatic Gla proteins than phylloquinone. The intake of menaquinones accounts for up to 25% of the total intake of vitamin K, and should there be a higher bioavailability, menaquinones would be important to consider in their contribution to human health. Indeed, certain observational studies have indicated that high intakes of menaquinones may be associated with greater reductions of vascular calcification and the risk of CVD than comparable amounts of phylloquinone.
Such effects have not been observed for bone health. However, these data are limited to observational studies conducted among Dutch or Japanese populations. Moreover, food composition tables for menaquinones are limited and available only for a few countries. In addition, studies investigating the bioavailability of menaquinones using stable isotope techniques are lacking. Therefore, research is warranted to compile more elaborate food composition data of menaquinones and more accurate data on the intake of menaquinones, at different stages of the life cycle (Table 3). These data should be used to investigate the relationship with disease incidence in populations other than the Dutch or Japanese. Stable isotope studies are required to quantify differences in absorption, bioavailability and distribution over the body between individual menaquinone forms and phylloquinone. Finally, intervention studies with clinical endpoints and a more elaborate validation of biomarkers for vitamin K status are required to quantify how the bioavailability and tissue distribution of menaquinones affect vitamin K status and health outcomes.

Clearly, significant gaps in the current knowledge on menaquinones exist. However, there is merit for considering both menaquinones and phylloquinone when developing future recommendations for vitamin K intake.

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