Effect of vitamin E intake from food and supplement sources on plasma α- and γ-tocopherol concentrations in a healthy Irish adult population

Yang Zhao, Frank J. Monahan, Breige A. McNulty, Mike J. Gibney and Eileen R. Gibney*
UCD Institute of Food and Health, University College Dublin, Dublin 4, Republic of Ireland

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Abstract
Vitamin E is believed to play a preventive role in diseases associated with oxidative stress. The aims of the present study were to quantify vitamin E intake levels and plasma concentrations and to assess dietary vitamin E adequacy in Irish adults. Intake data from the National Adult Nutrition Survey were used; plasma samples were obtained from a representative cohort of survey participants. Plasma α- and γ-tocopherol concentrations were measured by HPLC. The main sources of vitamin E in the diet were ‘butter, spreadable fats and oils’ and ‘vegetables and vegetable dishes’. When vitamin E intake from supplements was taken into account, supplements were found to be the main contributor, making a contribution of 29.2% to vitamin E intake in the total population. Supplement consumers had significantly higher plasma α-tocopherol concentrations and lower plasma γ-tocopherol concentrations when compared with non-consumers. Consumers of vitamin E supplements had significantly higher vitamin E intake levels and plasma α-tocopherol concentrations compared with consumers of other types of supplements, such as multivitamin and fish oil. Comparison with the Institute of Medicine Estimated Average Requirement of 12 mg/d indicated that when vitamin E intake from food and supplement sources was taken into account, 100% of the study participants achieved the recommended intake levels. When vitamin E intake from food sources was taken into account, only 68.4% of the females were found to achieve the recommended intake levels compared with 99.2% of the males. The results of the present study show that dietary vitamin E intake has a significant effect on plasma α- and γ-tocopherol concentrations. Furthermore, they show that the consumption of supplements is a major contributor to overall intake and has a significant effect on plasma vitamin E concentrations in the Irish population.

Key words: Vitamin E; Supplement types; Plasma tocopherol concentrations; Dietary intake

Vitamin E is an important lipid-soluble antioxidant in vivo and is essential for the normal functioning of the musculoskeletal, reproductive, neural and vascular systems(1). Vitamin E in the membrane of cells protects membranal phospholipid PUFA from oxidation, mediated by lipid radicals produced in the lipid peroxidation chain reaction(2). Vitamin E is believed to play a preventive role in diseases associated with oxidative stress such as cancer, CVD, cataract formation, age-related macular degeneration and diabetes mellitus(3–6).

Vitamin E exists as eight different isomers, four tocopherols (α-, β-, γ- and δ-tocopherols) and four corresponding tocotrienols(7). α-Tocotrienol is reported to have the highest biological activity, with the body preferentially absorbing and using this form(8). In contrast to α-tocopherol, γ-tocopherol is a powerful nucleophile that traps electrophilic mutagens in lipophilic compartments, which may protect lipids, DNA and proteins from peroxynitrite-dependent damage(9). The predominant form of vitamin E in food sources is γ-tocopherol, which is found in vegetable oils, grains, nuts and seeds(10). A survey carried out in the USA estimated that approximately 70% of the vitamin E intake from food sources occurs in the form of γ-tocopherol(10). The common form of vitamin E used in supplements is the acetate or succinate ester of α-tocopherol(11).

Studies have found that supplementation is generally related to increased dietary intake and plasma status of nutrients(12,13). In the Danish National Survey of Dietary Habits and Physical Activity (2000–4), supplement consumers were found to have a significantly higher intake of most micronutrients compared with non-consumers(12), and in the Canadian Health Measures Survey, the consumption of vitamin D supplements was found to contribute to a higher plasma status of 25-hydroxyvitamin D(13). Some surveys have also reported a positive correlation between vitamin E intake and plasma concentrations of tocopherols(14,15). A French observational study showed that dietary vitamin E intake had a significant influence on plasma α-tocopherol concentrations(16). In the US Women’s Health Initiative 1994–6,

Abbreviations: EAR, estimated average requirements; EU, European Union; IOM, Institute of Medicine; supplements, vitamin E-containing supplements.

* Corresponding author: Dr E. R. Gibney, email eileen.gibney@ucd.ie
it has been reported that vitamin E intake from supplements has a greater influence on plasma α- and γ-tocopherol concentrations than that from dietary sources. In addition, supplemental vitamin E intake was found to be the strongest predictor of high plasma α-tocopherol concentrations and low plasma γ-tocopherol concentrations in the US study. α-Tocopherol supplements may reduce circulating γ-tocopherol concentrations due to competition for hepatic transfer. In many surveys, vitamin E-containing supplements (hereafter referred to as supplements) were included in the estimation of daily dietary intake and their relative contributions to intake were not considered. Moreover, the effect of specific types of supplements on vitamin E intake levels and plasma α- and γ-tocopherol concentrations remains relatively unknown. Although some information is available on nutrient intake from food and supplement sources in the Irish adult population, neither plasma α- and γ-tocopherol concentrations nor the association between vitamin E intake and plasma α- and γ-tocopherol concentrations has been studied.

The aims of the present study were to determine mean daily vitamin E intake levels, assess dietary vitamin E intake adequacy compared with the recommended intake levels, and investigate the effect of vitamin E intake from food and supplement sources on plasma α- and γ-tocopherol concentrations in the Irish adult population.

Experimental design

Study population

Data for the present analysis were derived from the National Adult Nutrition Survey (NANS), a cross-sectional dietary survey carried out in 1500 healthy adults (740 males and 760 females) aged ≥18 years from across the Republic of Ireland during the years 2008–2010. Ethical approval for the survey was obtained from the Human Ethics Research Committee of University College Dublin and the University College Cork Clinical Research Ethics Committee of the Cork Teaching Hospitals. Individuals who agreed to participate in the survey signed a written consent form in accordance with the Declaration of Helsinki. The participants were representative of the Irish adult population in terms of age, sex, social class and urban/rural distribution. More detailed descriptions of the sampling procedures and the survey methodology have been published elsewhere. Of the 1500 participants, 1129 provided a blood sample, and data obtained from these subjects form the basis for the present study.

Collection of dietary intake data

Food and beverage intake data were collected using a 4-d semi-weighed food record, as described elsewhere. The participants recorded detailed information on the amount and type of all foods, beverages and nutritional supplements consumed over four consecutive days and, where applicable, recorded details regarding recipes, cooking methods and leftovers. The participants recorded their food intake at the brand level where possible and kept aside the packaging of foods that they consumed, which was later used to update the Irish National Food and Ingredient Database version 3.0. A quantification protocol established by the Irish Universities Nutrition Alliance for the North/South Ireland Food Consumption Survey in 1997–9 was updated for the NANS and used during data entry. Food intake was evaluated using Weighed Intake Software Package (WISP) version 3.0 (Tinuviel Software), which used data from McCance and Widdowson’s ‘The Composition of Food’ fifth and sixth editions, plus all the nine supplemental volumes. Each food and beverage was allocated an individual food code and subsequently assigned to one of the thirteen food groups for ease of analysis. Modifications were made to the food composition database to include recipes of composite dishes, nutritional supplements, generic Irish foods that are commonly consumed and new foods on the market.

Estimation of vitamin E intake

WISP provides information on the α-tocopherol equivalents (total vitamin E intake) from foods, but does not distinguish between the natural form of vitamin E and the synthetic forms used in supplements. Therefore, to estimate vitamin E intake from both food sources and supplement sources, supplements were first identified based on the presence of the vitamin in the ingredient list using the Irish National Food and Ingredient Database version 3.0 or were checked directly in the retail outlet or on the manufacturers’ website. Second, existing foods in the database were updated to reflect the current levels of α-tocopherol equivalents by comparing with the US Department of Agriculture-National Nutrient Database for Standard Reference. Overall, three nutrient descriptors were created for vitamin E intake, and these will be referred to throughout the article as follows: (1) all sources – vitamin E naturally occurring in foods and vitamin E from supplements; (2) food sources – vitamin E naturally occurring in only foods; (3) supplement sources – vitamin E from supplements. Vitamin E intake from all sources, food sources and supplement sources was adjusted for energy intake (mg/10 MJ) in the total population. Among the nineteen food groups, seven individually contributed less than 3.7% to vitamin E intake and so were combined into one food group termed ‘Others’. In total, thirteen food groups were taken into consideration as all sources of vitamin E intake.

A common qualitative classification of supplements was developed. The categories were ‘multivitamin’, ‘fish oil’, ‘seed oil’, ‘fish and seed oil’, and ‘vitamin E’. A category termed ‘Others’ comprised non-specified vitamin E-containing supplements, including protein powders, protein and energy drinks, weight-loss products, isoflavone supplements, Se supplements and Mg supplements. Supplement consumers who consumed more than one type of supplement (multi-supplement consumers) were categorised using two approaches. First, as the number of supplement consumers of both multivitamin supplements and fish oil was greater than that of consumers of other supplement combinations, they were assigned to the category ‘multivitamin'.
and fish oil'. Second, the remaining multi-supplement consumers \((n = 21)\) were assigned to the category of supplementation with the greatest contribution to vitamin E intake from supplement sources. Therefore, each supplement consumer belonged to only one category: ‘multivitamin’, ‘fish oil’, ‘seed oil’, ‘fish and seed oil’, ‘vitamin E or ‘Others’.

### Blood sampling and biochemical analysis

The blood collection protocol used in the NANS has been described elsewhere\(^{3,36}\). Blood processing and sample fractions were performed at biological laboratories in University College Dublin and University College Cork, and samples were stored at \(-80^\circ C\) until analysis. All plasma samples for vitamin E analysis were collected in an EDTA-containing tube. Plasma vitamin E (\(\alpha-\) and \(\gamma-\)tocopherol) concentrations were determined by reversed-phase HPLC according to the method of Siluk \textit{et al.} with slight modifications\(^{37}\). Aliquots (200 \(\mu\)l) of plasma were vortex-mixed with 200 \(\mu\)l of distilled water, 400 \(\mu\)l of ethanol (containing 0.04 \% butylated hydroxytoluene) and 20 \(\mu\)l of internal standard (15 \(\mu\)g/ml \(\alpha\)-tocopherol acetate in ethanol). Extraction of \(\alpha\)- and \(\gamma\)-tocopherols was performed with 800 \(\mu\)l hexane by vortex-mixing for 3 min and centrifuging at 5600 \(g\) for 10 min at 10\({^\circ}\)C. The hexane extract was collected, and the solvent was evaporated using Speed Vac (Genevac). The residue was dissolved in 200 \(\mu\)l of methanol, vortex-mixed for 30 s and transferred into an amber vial. The methanol extract (20 \(\mu\)l) was analysed on an Agilent 1200 series HPLC system (Agilent Technologies, Inc.). Separation was achieved on a Zorbax Eclipse XDB-C18 column (150 mm \(\times\) 4 mm inner diameter; 5 \(\mu\)m particle size) with a RX-C8 guard column (12.5 mm \(\times\) 4.6 mm inner diameter; 5 \(\mu\)m particle size). The column was attached to a 1200 series diode array detector (Agilent Technologies, Inc.), set at an excitation wavelength of 295 nm for \(\alpha\)- and \(\gamma\)-tocopherols and of 298 nm for \(\alpha\)-tocopherol acetate\(^{37}\). The mobile phase was 100 \% methanol at a flow rate 1 ml/min.

A UV spectrophotometer (UV mini-1240; Shimadzu) was used to determine the concentrations of standard solutions of \(\alpha\)-tocopherol, \(\gamma\)-tocopherol and \(\alpha\)-tocopherol acetate according to published molar extinction coefficients\(^{38}\). Plasma \(\alpha\)- and \(\gamma\)-tocopherol concentrations were calculated by comparing with a corresponding calibration curve. The recovery of the internal standard was 98 \% for \(\alpha\)-tocopherol and 101 \% for \(\gamma\)-tocopherol. This is similar to original findings from the method of Siluk \textit{et al.}\(^{37}\), where the accuracy of recovery ranged from 96 to 106 \% for \(\alpha\)-tocopherol and from 101 to 109 \% for \(\gamma\)-tocopherol.

### Statistical analyses

All statistical analyses were carried out using SPSS version 20.0 (SPSS, Inc.). Compliance to the current RDA\(^{39}\) and estimated average requirements (EAR)\(^{40}\) was estimated using the method of Wearne & Day\(^{41}\). Of the 1129 participants, 420 who were considered to have under-reported energy intake using the Goldberg method\(^{42}\) were excluded from the compliance analysis, leaving 709 participants. All the 1129 participants were included in all other analyses. Vitamin E intake from all sources, food sources and supplement sources was calculated, and the relative contributions of the thirteen food groups to vitamin E intake in the total population were evaluated. Energy-adjusted vitamin E intake (mg/10 MJ) from all sources, food sources and supplement sources was calculated for the total population, and differences across age groups (18–35, 36–50, 50–64 and >64 years) and sexes were determined using a two-way ANOVA, followed by Tukey’s post hoc test. Data on the intake levels and plasma concentrations of vitamin E were skewed and therefore log-transformed before statistical analyses. Data on plasma \(\alpha\)- and \(\gamma\)-tocopherol concentrations of the total population and supplement consumers and non-consumers within each age group and sex were analysed using two-way ANOVA. The participants were also divided into quartiles of vitamin E intake from all sources, food sources and supplement sources, and a general linear model univariate analysis (adjusted for age and sex) was used to assess differences in plasma \(\alpha\)- and \(\gamma\)-tocopherol concentrations across quartiles of intake.

### Results

#### Study population characteristics

A total of 1129 Irish adults were included in the present study. The participants were representative of the Irish adult population in terms of age, sex, social class and location\(^{18,20}\). In total, 70 \% of the males and 56 \% of the females were classified as overweight/obese, and 20 \% of the participants were current smokers. Furthermore, 31 \% of the participants consumed at least one nutritional supplement, with more females (36 \%) consuming nutritional supplements than males (26 \%) (online supplementary Table S1).

#### Vitamin E intake in the total population

The mean daily vitamin E intake from all sources, food sources and supplement sources was 13·8, 9·8 and 21·9 mg/d, respectively, in the total population (online supplementary Table S2). There was a stepwise increase in vitamin E intake from all sources across age groups, with the lowest intake being found in the 18–35 years age group. Vitamin E intake from all sources was slightly higher in females than in males (14·5 \(v.\) 13·2 mg/d). However, this seemed to be due to supplement intake, as female supplement consumers had a higher intake than male supplement consumers (28·6 \(v.\) 14·4 mg/d). Vitamin E intake from food sources was higher in males than in females (10·7 \(v.\) 8·9 mg/d).

The percentage of Irish adults meeting dietary vitamin E intake requirements, including the European Union (EU) RDA (12 mg/d)\(^{39}\) and the Institute of Medicine (IOM) EAR\(^{40}\), was determined (Table 1). On using the IOM EAR or EU RDA of 12 mg/d, 100 \% of the study population was found to achieve the recommended intake levels; however, when intake from food sources was considered, only 99 \% of the males and 68 \% of the females were found to achieve the recommended intake levels. This disparity between
males and females is probably due to the absence of sex-specific recommendations, with an intake of 12 mg/d being recommended for both males and females; as such males, with considerably higher energy and fat intakes than females, more readily achieve this recommended intake level \(^{(20)}\). Less than 1 % of the participants were found to have intake levels above the tolerable upper intake level of 300 mg/d \(^{(43)}\) (data not shown).

The percentage of contribution of the thirteen food groups to the mean daily vitamin E intake in the total population was assessed (Fig. 1). When vitamin E intake from food sources was taken into account, the food group ‘butter, spreadable fats and oils’ was found to be the main source of dietary vitamin E intake, followed by ‘vegetables and vegetable dishes’. When vitamin E intake from supplements was taken into account, supplements were found to make a significant contribution, accounting for 29 % of dietary vitamin E intake in the total population. Of the 31 % of participants taking nutritional supplements, 18 % (17 % of the males and 20 % of the females) consumed at least one supplement specifically containing vitamin E. When supplement consumers and non-consumers were analysed separately, ranking of the contribution of food groups to vitamin E intake remained the same, across both sets of participants. Thus, supplement consumers and non-consumers seemed to have similar dietary patterns with respect to vitamin E intake.

The energy-adjusted mean daily vitamin E intake (mg/10 MJ per d) from all sources, food sources and supplement sources is summarised in Table 2. A stepwise increase in vitamin E intake from all sources and food sources was observed in the total population with an increase in age \((P<0.001)\). A significant effect of sex was found across quartiles of intake from all sources, food sources and supplement sources, with a higher vitamin E intake being found in females than in males \((P<0.001)\). No age \(\times\) sex interactions were observed across groups.

### Plasma vitamin E concentrations in the Irish population

Plasma \(\alpha\)- and \(\gamma\)-tocopherol concentrations of the total population, supplement consumers and non-consumers are given in Tables 3 and 4, respectively. Supplement consumers had significantly higher plasma \(\alpha\)-tocopherol concentrations \((P<0.001)\) and lower plasma \(\gamma\)-tocopherol concentrations \((P<0.001)\) when compared with non-consumers across all age groups. The mean plasma \(\alpha\)- and \(\gamma\)-tocopherol concentrations of supplement consumers, across all age groups, were 28.1 and 1.5 \(\mu\)mol/l, while those of non-consumers

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**Table 1. Percentage of Irish adults meeting dietary vitamin E intake requirements including the Institute of Medicine estimated average requirement (12 mg/d)**

<table>
<thead>
<tr>
<th></th>
<th>Total population</th>
<th>Males</th>
<th>Females</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>All sources</strong></td>
<td>100·0</td>
<td>100·0</td>
<td>100·0</td>
</tr>
<tr>
<td><strong>Food sources</strong></td>
<td>86·9</td>
<td>99·2</td>
<td>68·4</td>
</tr>
</tbody>
</table>

*Includes intake from both food and supplement sources.*
Effect of age and sex on plasma vitamin E concentrations

There were significant differences in plasma α-tocopherol concentrations among the age groups (P<0.001) in the total population, non-consumers and supplement consumers, with the concentrations increasing with an increase in age (Table 3). An age × sex interaction was observed in the total population, with males in the 51–64 and >64 years age groups, but not in the 18–35 and 36–50 years age groups, having lower plasma α-tocopherol concentrations compared with females in the same age groups (P=0.007; Table 3).

When plasma γ-tocopherol concentrations were evaluated (Table 4), an effect of age was found in the total population (P=0.004) and non-consumers of supplements (P=0.001), whereby the 51–64 years age group had higher plasma γ-tocopherol concentrations than the other age groups. No sex effect or age × sex interactions were observed across groups.

Effect of food and supplement sources on plasma vitamin E concentrations

To further investigate the effect of vitamin E intake sources on plasma α- and γ-tocopherol concentrations, the total population was divided into quartiles of vitamin E intake from all sources (Table 5) and food sources (Table 6), while supplement consumers were divided into quartiles of vitamin E intake from supplement sources (Table 7). Plasma α- and γ-tocopherol concentrations were significantly different across quartiles of vitamin E intake from all sources.

Table 2. Effect of age and sex on the energy-adjusted mean daily vitamin E intake (mg/10 MJ per d) from all sources, food sources and supplement sources (Number of participants, mean values and standard deviations)

<table>
<thead>
<tr>
<th>Age groups (years)</th>
<th>18–35</th>
<th>36–50</th>
<th>51–64</th>
<th>&gt;64</th>
<th>P (two-way ANOVA)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>All sources</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>Age Sex Age × sex</td>
</tr>
<tr>
<td>All males</td>
<td>428 ± 13.6 ± 29.4</td>
<td>332 ± 15.1 ± 28.0</td>
<td>224 ± 20.0 ± 37.4</td>
<td>145 ± 22.2 ± 54.0</td>
<td>0.001</td>
</tr>
<tr>
<td>Females</td>
<td>196 ± 13.6 ± 11.8</td>
<td>154 ± 11.5 ± 6.4</td>
<td>114 ± 13.4 ± 9.0</td>
<td>66 ± 13.5 ± 6.8</td>
<td>0.001</td>
</tr>
<tr>
<td>Food sources</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>Age Sex Age × sex</td>
</tr>
<tr>
<td>All males</td>
<td>428 ± 10.4 ± 4.2</td>
<td>332 ± 11.4 ± 4.4</td>
<td>224 ± 12.6 ± 5.6</td>
<td>145 ± 13.2 ± 6.3</td>
<td>0.001</td>
</tr>
<tr>
<td>Females</td>
<td>196 ± 11.2 ± 4.3</td>
<td>178 ± 13.7 ± 4.2</td>
<td>110 ± 26.7 ± 5.6</td>
<td>79 ± 29.6 ± 7.2</td>
<td>0.001</td>
</tr>
<tr>
<td>Supplement sources</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>Age Sex Age × sex</td>
</tr>
<tr>
<td>All males</td>
<td>65 ± 21.2 ± 59.5</td>
<td>49 ± 25.6 ± 67.4</td>
<td>59 ± 28.1 ± 67.9</td>
<td>35 ± 37.6 ± 103.4</td>
<td>0.001</td>
</tr>
<tr>
<td>Females</td>
<td>36 ± 25.3 ± 77.2</td>
<td>22 ± 6.7 ± 9.9</td>
<td>25 ± 8.7 ± 11.2</td>
<td>15 ± 4.7 ± 4.5</td>
<td>0.001</td>
</tr>
</tbody>
</table>

Table 3. Effect of age and sex on plasma α-tocopherol concentrations (µmol/l) in the total population, supplement consumers and non-consumers (Number of participants, mean values and standard deviations)

<table>
<thead>
<tr>
<th>Age groups (years)</th>
<th>18–35</th>
<th>36–50</th>
<th>51–64</th>
<th>&gt;64</th>
<th>P (two-way ANOVA)†</th>
</tr>
</thead>
<tbody>
<tr>
<td>All population</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>Age Sex Age × sex</td>
</tr>
<tr>
<td>All males</td>
<td>428 ± 22.18 ± 5.30</td>
<td>332 ± 27.22 ± 6.81</td>
<td>224 ± 28.62 ± 7.24</td>
<td>145 ± 27.59 ± 7.78</td>
<td>0.001</td>
</tr>
<tr>
<td>Females</td>
<td>196 ± 22.16 ± 4.97</td>
<td>178 ± 26.98 ± 6.38</td>
<td>110 ± 29.52 ± 6.50</td>
<td>79 ± 29.15 ± 7.92</td>
<td>0.001</td>
</tr>
<tr>
<td>Non-consumers</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>Age Sex Age × sex</td>
</tr>
<tr>
<td>All males</td>
<td>363 ± 21.83 ± 4.81</td>
<td>283 ± 26.73 ± 6.48</td>
<td>165 ± 28.37 ± 7.14</td>
<td>110 ± 26.67 ± 7.43</td>
<td>0.001</td>
</tr>
<tr>
<td>Females</td>
<td>196 ± 22.02 ± 4.99</td>
<td>132 ± 27.15 ± 7.40</td>
<td>89 ± 28.01 ± 8.24</td>
<td>51 ± 25.39 ± 6.97</td>
<td>0.001</td>
</tr>
<tr>
<td>Supplement consumers</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>n Mean ± SD</td>
<td>Age Sex Age × sex</td>
</tr>
<tr>
<td>All males</td>
<td>65 ± 24.12 ± 7.21</td>
<td>49 ± 30.09 ± 7.97</td>
<td>59 ± 29.32 ± 7.52</td>
<td>35 ± 30.48 ± 8.24</td>
<td>0.001</td>
</tr>
<tr>
<td>Females</td>
<td>36 ± 23.10 ± 8.07</td>
<td>22 ± 29.60 ± 6.31</td>
<td>25 ± 26.80 ± 6.14</td>
<td>15 ± 26.85 ± 8.18</td>
<td>0.001</td>
</tr>
</tbody>
</table>

*Mean values were significantly different between sexes within each age group (P<0.05).
†Differences across age groups and sexes were assessed using two-way ANOVA followed by Tukey’s post hoc test.
(P<0.001 and 0.005, respectively), with plasma α-tocopherol concentrations increasing (by 9%) and plasma γ-tocopherol concentrations decreasing (by 13%) with an increase in intake. There was a significant effect of food sources on plasma α-tocopherol concentrations (P=0.027), which increased by 6%, but there was no change in plasma γ-tocopherol concentrations across quartiles of intake (Table 6). Similarly, in supplement consumers, a significant effect of vitamin E intake from supplement sources on plasma α-tocopherol concentrations (P=0.001), but not on plasma γ-tocopherol concentrations, was observed across quartiles of intake. However, among supplement consumers, those in the highest quartile had significantly higher intake levels (72 mg/d) and higher plasma α-tocopherol concentrations (32 μmol/l).

The range of manufacturers’ recommended doses in the different types of supplements was large, with one ‘vitamin E’ supplement providing a high dose (546 mg/d), while the lowest recommended dose (0.2 mg/d) was observed in fish oil-type supplements (Table 8). The vast majority of supplement consumers consumed either fish oil or multi-vitamin supplements, with only 0.4% of the total population consuming either fish oil or multi-vitamin supplements (Table 8). The vast majority of supplement consumers consumed fish oil-type supplements (Table 8).

Discussion

In the present study, Irish adults were found to generally achieve the EU RDA and IOM EAR of 12 mg/d. The raw vitamin E intake values reported herein (13 and 15 mg/d from all sources and 11 and 9 mg/d from food sources for males and females, respectively) are comparable to those reported in other international studies (44–47). The Polish National Nutrition Intake Survey (2001) reported mean daily vitamin E intake values, including for supplement sources, of 16 mg/d for males and 11 mg/d for females (46). The UK National Diet and Nutrition Survey (NDNS, 2004) published values of 13 and 15 mg/d for the mean daily vitamin E intake from all sources of different types of supplements (P<0.001), with ‘vitamin E’ supplements providing a considerably higher amount of vitamin E than all other types of supplements. No significant difference was observed in vitamin E intake from food sources across consumers of different types of supplements, but there was a significant difference in plasma α-tocopherol concentrations (P<0.001) across consumers of different types of supplements, with ‘vitamin E’ supplement consumers having significantly higher plasma α-tocopherol concentrations than consumers of other types of supplements.

Table 5. Plasma α- and γ-tocopherol concentrations (μmol/l) across quartiles of daily vitamin E intake from all sources

<table>
<thead>
<tr>
<th>Quartile</th>
<th>Intake levels</th>
<th>α-Tocopherol concentrations</th>
<th>γ-Tocopherol concentrations</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4.92b</td>
<td>24.68b</td>
<td>1.81a</td>
</tr>
<tr>
<td>2</td>
<td>8.08b</td>
<td>25.29b</td>
<td>1.76b</td>
</tr>
<tr>
<td>3</td>
<td>11.39b</td>
<td>25.59b</td>
<td>1.75b</td>
</tr>
<tr>
<td>4</td>
<td>31.00b</td>
<td>26.98b</td>
<td>1.57b</td>
</tr>
</tbody>
</table>

GLM, general linear model.

P (GLM)*

<0.001

<0.001

<0.001

<0.001

<0.001

<0.001

<0.001

<0.001

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<0.001
suggested that the consumption of supplements by the Irish population has increased over the last 10 years. The UK survey (NDNS, 2004) also demonstrated the considerable contribution of supplement sources to vitamin E intake in the total population, increasing the mean daily intake over that from food sources by 26 % for males and 85 % for females. In contrast to the present results, in the USA (NHANES 2003–6), 58 % of adults were found to have intake levels below the IOM EAR when intake from supplements was taken into account.

Comparison of the mean daily intake values of the study population with the IOM EAR or EU RDA showed that the prevalence of inadequate vitamin E intake from food sources was less than 1 % in males and 32 % in females; however, 100 % of both males and females achieved the recommended intake levels when intake from food and supplement sources was taken into account, indicating that the consumption of supplements contributed to dietary adequacy in females. The reason for 99 % of the Irish males achieving the recommended intake from food sources alone is that they had significantly higher energy and fat intakes compared with females, which contributed to a higher vitamin E intake. Furthermore, in the present study, the consumption of supplements was found to contribute 29 % to vitamin E intake in the total population; a previous Irish national consumption survey published 10 years ago (The North/South Irish Food Consumption Survey) has reported that supplement use contributed only 6 % to the mean daily vitamin E intake in males and 12 % in females. This suggests that the consumption of supplements by the Irish population has increased over the last 10 years. The UK survey (NDNS, 2004) also demonstrated the considerable contribution of supplement sources to vitamin E intake in the total population, increasing the mean daily intake over that from food sources by 26 % for males and 85 % for females. In contrast to the present results, in the USA (NHANES 2003–6), 58 % of adults were found to have intake levels below the IOM EAR when intake from supplements was taken into account.

The main sources of vitamin E in the Irish adult diet are ‘butter, spreadable fats and oils’ and ‘vegetables and vegetable dishes’. ‘Bread, spreadable fats and oils’ are rich sources of vitamin E and also consumed very frequently in small portions by Irish adults, with an average of 4–4 % energy being derived from fat in this cohort. ‘Vegetables and vegetable dishes’ are consumed very frequently in large portions (148 g/d) and vegetable dishes often contain added butter and sauce, which are rich sources of vitamin E. In agreement with the present findings, the main contributor to dietary vitamin E intake in the UK survey (NDNS, 2004) was also ‘fat spreads’, making a contribution of 18 % to dietary vitamin E intake; however, the second contributor was ‘cereals and cereal products’, making a contribution of 17 % to dietary vitamin E intake. It is important to note that ‘breakfast cereals’, which were categorised within ‘cereals and cereal products’ in the UK survey, contributed 5 % to dietary intake in the total population. This is similar to the present study where the ‘breakfast cereals’ group contributed 3·8 % to Irish vitamin E intake from all sources. The US Continuing Survey of Food Intakes by Individuals (CSFII) carried out from 1994 to 1996 reported that the two main contributors to dietary vitamin E from food sources and of 11 and 8 mg/d for that from food sources for males and females, respectively, and the Survey of Lifestyle, Attitudes and Nutrition in Ireland (SLAN 2007) reported values of 8·8 mg/d and 9·1 mg/d for the mean dietary vitamin E intake from food sources for males and females, respectively. The US National Health and Nutrition Examination Survey (NHANES) 2001–2 reported values of 7·0 and 6·1 mg/d for the mean vitamin E intake from food sources for males and females, respectively.

### Table 6. Plasma α- and γ-tocopherol concentrations (µmol/l) across quartiles of daily vitamin E intake from food sources

(Mean values and standard deviations, n 1129)

<table>
<thead>
<tr>
<th>Quartiles</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>P (GLM)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>Intake levels</td>
<td>4·66d</td>
<td>1·10</td>
<td>7·62c</td>
<td>0·72</td>
<td>10·38b</td>
</tr>
<tr>
<td>α-Tocopherol concentrations</td>
<td>24·79b</td>
<td>7·03</td>
<td>25·70ab</td>
<td>6·26</td>
<td>25·73ab</td>
</tr>
<tr>
<td>γ-Tocopherol concentrations</td>
<td>1·77</td>
<td>0·87</td>
<td>1·78</td>
<td>0·92</td>
<td>1·64</td>
</tr>
</tbody>
</table>

GLM, general linear model.

Mean values with unlike superscript letters were significantly different between the quartiles (P<0·05).

* GLM univariate analysis, adjusted for age and sex, was carried out to determine significance.

### Table 7. Plasma α- and γ-tocopherol concentrations (µmol/l) across quartiles of daily vitamin E intake from supplement sources (only consumers)

(Mean values and standard deviations, n 208)

<table>
<thead>
<tr>
<th>Quartiles</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>P (GLM)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
<td>SD</td>
<td>Mean</td>
</tr>
<tr>
<td>Intake levels</td>
<td>0·51b</td>
<td>0·47</td>
<td>4·84b</td>
<td>1·96</td>
<td>10·09b</td>
</tr>
<tr>
<td>α-Tocopherol concentrations</td>
<td>26·13b</td>
<td>8·26</td>
<td>27·34b</td>
<td>6·80</td>
<td>27·27b</td>
</tr>
<tr>
<td>γ-Tocopherol concentrations</td>
<td>1·58</td>
<td>0·74</td>
<td>1·68</td>
<td>1·08</td>
<td>1·36</td>
</tr>
</tbody>
</table>

GLM, general linear model.

Mean values with unlike superscript letters were significantly different between the quartiles (P<0·05).

* GLM univariate analysis, adjusted for age and sex, was carried out to determine significance.
intake in the US diet are 'ready-to-eat cereals' and 'cakes, cookies, pies and doughnuts'. In the US survey, the 'ready-to-eat cereals' group contained products fortified with synthetic vitamin E, which is not the case in Ireland or the UK. In the present study, only 6% of the ready-to-eat cereals consumed were fortified with vitamin E. It is also worth noting that synthetic forms of vitamin E are less bioavailable than the natural form, and Maras et al. suggested that the actual available dietary vitamin E amounts from 'ready-to-eat cereals' may be lower than those reported in the US publication for this reason. Similar to a previous study, in the present study also, it was found that 0.7% of the foods (eighteen from a total of 2552 foods) were fortified with vitamin E and that fortification with vitamin E had little impact on total vitamin E intake. Thus, in the present study, foods fortified with vitamin E were not selected as an individual food group due to very low numbers. To our knowledge, the effect of fortified foods on plasma vitamin E concentrations at a population level has not been investigated.

The effects of age and sex on vitamin E intake were clearly observed in the present study. Older participants had a higher vitamin E intake as they consumed more vitamin E-rich foods, such as dairy spreads, compared with younger participants (data not shown). A greater prevalence of supplement use was also found in those aged >64 years, particularly in females. The consumption of supplements by females was the main factor contributing to the sex effects. Among supplement consumers, vitamin E intake from supplements in females aged 36–50 years was five times higher than that in male supplement consumers, while vitamin E intake from supplements increased with an increase in age and that females consumed significantly more vitamin E from supplement sources than males. An effect of sex was also observed in the French Supplementation en Vitamines et Minéraux AntiOxydants (SU.VI.MAX) Study; the vitamin E intake levels of females were significantly higher than those of males.

An effect of age on plasma α-tocopherol concentrations was observed across the total population and in both supplement consumers and non-consumers, with plasma α-tocopherol concentrations increasing with age. This was in agreement with findings from a Swiss survey of non-consumers of supplements, where both plasma α- and γ-tocopherol concentrations were found to increase with age. The effect of sex on plasma α-tocopherol concentrations in the total population was attributable to the higher plasma α-tocopherol concentrations of female supplement consumers compared with their male counterparts, influenced by the amount of supplements consumed. Other studies have also suggested that the main indicator of plasma α- and γ-tocopherol concentrations is vitamin E supplement intake, where the plasma α-tocopherol concentrations of vitamin E supplement consumers were found to be substantially higher than those of non-consumers.

With regard to the effect of supplement sources on plasma α-tocopherol concentrations, only high vitamin E intake, as demonstrated by those in the fourth quartile of intake in the present study, had a significant effect. It is perhaps unsurprising that food sources are unlikely to provide such high vitamin E amounts and that supplement sources therefore have a larger impact on plasma α- and γ-tocopherol concentrations than food sources. A similar study in a Costa Rican population examined plasma α-tocopherol concentrations across the reported α-tocopherol intake quintiles and also found a
significant increase in plasma α-tocopherol concentrations across quintiles of intake, reporting that the significance was mainly attributable to those in the highest intake quintile. Similar to the present results, several studies have indicated that with increasing total vitamin E intake, plasma α-tocopherol concentrations increase and γ-tocopherol concentrations decrease (59,60), which is hypothesised to be due to competition for the α-tocopherol hepatic transfer protein. Studies have shown that the hepatic α-tocopherol transfer protein preferentially incorporates α-tocopherol into VLDL, potentially explaining the rise in α-tocopherol concentrations with increasing intakes and concomitant decrease in γ-tocopherol concentrations (59). Examination of the impact of factors such as smoking habits (16) and BMI (60), which are known to have an effect on plasma α- and γ-tocopherol concentrations, was beyond the scope of the present study.

The present study is one of the first to examine the impact of supplement types on vitamin E intake and plasma α- and γ-tocopherol concentrations in a national population. The results indicated that the types of supplements consumed had a significant impact on the reported vitamin E intake, reflecting the wide range of recommended doses in different types of supplements. As there was no significant difference in vitamin E intake from food sources among consumers of different types of supplements, the differences in intake were mainly attributable to the consumption of supplements. Among the consumers of different types of supplements, only ‘vitamin E’ supplement consumers had a significantly higher dietary vitamin E intake compared with other groups. In agreement with vitamin E intake quartile results, ‘vitamin E’ supplement consumers had significantly higher plasma α-tocopherol concentrations and slightly lower plasma γ-tocopherol concentrations when compared with consumers of other supplements.

In conclusion, vitamin E intake levels have been characterised for the first time in an Irish adult population. In general, the mean daily vitamin E intake of Irish adults was in accordance with the levels recommended by the EU and IOM, but females appeared to be reliant on supplementation to achieve these recommended levels. In Irish adults, vitamin E intake from supplements varied across consumers of different types of supplements, while plasma α- and γ-tocopherol concentrations were found to increase and decrease, respectively, with an increase in vitamin E intake. ‘Vitamin E’ supplements seemed to be the factors contributing to high vitamin E intake levels and plasma α-tocopherol concentrations.

Supplementary material

To view supplementary material for this article, please visit
http://dx.doi.org/10.1017/S0007114514002438

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