



# Association between nutritional profiles of foods underlying Nutri-Score front-of-pack labels and mortality: EPIC cohort study in 10 European countries

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Additional material is published online only. To view please visit the journal online.

# Cite this as: *BMJ*2020;370:m3173

http://dx.doi.org/10.1136/bmj.m3173

Accepted: 3 August 2020

### **ABSTRACT**

## **OBJECTIVE**

To determine if the Food Standards Agency nutrient profiling system (FSAm-NPS), which grades the nutritional quality of food products and is used to derive the Nutri-Score front-of-packet label to

guide consumers towards healthier food choices, is associated with mortality.

#### DESIGN

Population based cohort study.

#### **SETTING**

European Prospective Investigation into Cancer and Nutrition (EPIC) cohort from 23 centres in 10 European countries.

#### **PARTICIPANTS**

521 324 adults; at recruitment, country specific and validated dietary questionnaires were used to assess their usual dietary intakes. A FSAm-NPS score was calculated for each food item per 100 g content of energy, sugars, saturated fatty acids, sodium, fibre, and protein, and of fruit, vegetables, legumes, and nuts. The FSAm-NPS dietary index was calculated for each participant as an energy weighted mean of the FSAm-NPS score of all foods consumed. The higher the score the lower the overall nutritional quality of the diet.

## MAIN OUTCOME MEASURE

Associations between the FSAm-NPS dietary index score and mortality, assessed using multivariable adjusted Cox proportional hazards regression models.

#### DECILIT

After exclusions, 501 594 adults (median follow-up 17.2 years, 8 162 730 person years) were included in the analyses. Those with a higher FSAm-NPS dietary index score (highest versus lowest fifth) showed an increased risk of all cause mortality (n=53 112 events from non-external causes; hazard ratio 1.07, 95% confidence interval 1.03 to 1.10, P<0.001 for trend) and mortality from cancer (1.08, 1.03 to 1.13, P<0.001 for trend) and diseases of the circulatory (1.04, 0.98 to 1.11, P=0.06 for trend), respiratory (1.39, 1.22 to 1.59, P<0.001), and digestive (1.22,

## WHAT IS ALREADY KNOWN ON THIS TOPIC

Helping consumers make healthier food choices is a major challenge for the prevention of non-communicable diseases and related deaths

The Food Standards Agency nutrient profiling system (FSAm-NPS), which grades the nutritional quality of food products based on 100 g content of energy, sugars, saturated fatty acids, sodium, fibre and protein, and of fruit, vegetables, legumes, and nuts, underlies the Nutri-Score

Nutri-Score is a simple nutrition label selected by several countries in Europe and considered at the European Union level as a candidate for enabling uniform food labelling systems

FSAm-NPS defined nutritional quality of foods has been studied in relation to health but not to mortality in French cohorts, and recently to cancer risk in the large multinational European Prospective Investigation into Cancer and Nutrition (EPIC) cohort; evidence in an international setting for other health outcomes and especially mortality is still needed

## **WHAT THIS STUDY ADDS**

This study used data from EPIC, a large cohort comprising 501 594 adults from 10 European countries (53 112 deaths), with diverse profiles and dietary patterns and showed that the consumption of foods with higher FSAm-NPS scores (lower nutritional quality) was associated with increased all cause and cause specific mortality

These results add support to the relevance of using the FSAm-NPS (and the derived Nutri-Score) to characterise healthier food choices as a basis for public health nutritional policies in Europe

This is important considering ongoing and future debates at the EU level on making food labelling systems uniform on the front of food product packaging

1.02 to 1.45, P=0.03 for trend) systems. The age standardised absolute rates for all cause mortality per 10 000 persons over 10 years were 760 (men=1237; women=563) for those in the highest fifth of the FSAm-NPS dietary index score and 661 (men=1008; women=518) for those in the lowest fifth.

#### CONCLUSIONS

In this large multinational European cohort, consuming foods with a higher FSAm-NPS score (lower nutritional quality) was associated with a higher mortality for all causes and for cancer and diseases of the circulatory, respiratory, and digestive systems, supporting the relevance of FSAm-NPS to characterise healthier food choices in the context of public health policies (eg, the Nutri-Score) for European populations. This is important considering ongoing discussions about the potential implementation of a unique nutrition labelling system at the European Union level.

#### Introduction

Poor nutrition is a major well known risk factor for non-communicable diseases, with an estimated 11 million deaths from such diseases attributed to unhealthy diets worldwide in 2017. Although it is well established that less sugars, saturated fats, salt, and energy and more dietary fibres or fruit and vegetables should be consumed for better health, putting these recommendations into practice remains an important challenge. Helping consumers make healthier food choices could therefore serve as one of the key strategies to prevent mortality from non-communicable diseases. A front-of-pack label providing user friendly information on the nutritional quality of food products has been identified as a possible solution to this problem.2 3 Such labels have the potential to help consumers choose food products with a better nutritional quality at the point of purchase, and, simultaneously, to incentivise food manufacturers to improve the nutritional quality of products, thus contributing to a healthier food environment. 4 5 The Nutri-Score labelling system, which uses five colours, 6 is considered promising in a broad international context. Nutri-Score classifies food products into five categories according to nutritional quality (from category A, indicating higher nutritional quality, to category E, indicating lower nutritional quality) assessed using the Food Standards Agency nutrient profiling system (FSAm-NPS), an adapted version of a nutrient profiling system (FSA-NPS) initially developed by the British Food Standards Agency.<sup>7-9</sup> This scoring system was developed to prevent a large range of nutrition related non-communicable diseases, by allocating a score to a given food or beverage per 100 g content of energy, saturated fatty acids, sugars, sodium, dietary fibre, and protein, and of fruit, vegetables, legumes, and nuts.

In 2017, public health authorities in France officially adopted Nutri-Score<sup>10</sup> <sup>11</sup> after a series of studies showed the validity, scientific relevance, and potential public health benefits of the FSAm-NPS<sup>12-14</sup> and of the Nutri-Score label as a tool for public health nutrition

policies<sup>15-23</sup> (reviewed in<sup>24</sup>). Subsequently, Belgium, Spain, Germany, the Netherlands, Switzerland, and Luxembourg adopted Nutri-Score. Medical professionals and academic societies in Europe have also recognised the importance and potential public health impact of Nutri-Score as a tool that can be recommended to the general public and patients to guide them towards food choices of higher nutritional quality.

Under current European Union labelling regulations, member states cannot legally enforce the inclusion of a front-of-pack nutrition label such as Nutri-Score, which leaves the choice to food manufacturers. The stakes are therefore high for standardised nutritional labelling systems at the EU level, using a unique mandatory front-of-pack nutrition label. Similar discussions are ongoing in America and Australia. Since most of the original studies assessing the validity of the FSAm-NPS underlying Nutri-Score were performed in France, 24-27 it is important that the validity of the model is extended to international settings to provide relevant scientific evidence for ongoing discussions in the EU and beyond.

Part of the validity assessment of FSAm-NPS is to study the association between the nutritional quality of food products graded by the scoring system and health outcomes. Such studies, done in the French SU.VI.MAX and NutriNet-Santé cohorts, showed that on average consumption of more food products with lower FSAm-NPS scores (representing higher nutritional quality) was associated with more favourable outcomes for weight gain,<sup>29</sup> asthma symptoms,<sup>27</sup> metabolic syndrome,<sup>30</sup> cardiovascular diseases. 31 32 and cancer. 33 34 Recently. we showed that similar observations could be made for cancer risk in a large multinational European cohort, the European Prospective Investigation into Cancer and Nutrition (EPIC) study.35 In the current study we investigated the association between the FSAm-NPS scores of food products consumed and mortality in this large and diverse European population.

# Methods

## Study population: EPIC cohort

This study was conducted within the framework of the EPIC cohort study (https://epic.iarc.fr/), which enrolled more than 500 000 volunteers (aged 25-70 years) from 23 centres in 10 European countries (Denmark, France, Germany, Greece, Italy, Netherlands, Norway, Spain, Sweden, and the United Kingdom) between 1992 and 2000. This cohort study investigates metabolic, dietary, lifestyle, and environmental factors associated with the development of cancer and other noncommunicable diseases in Europe. All participants gave written informed consent. Details of the study design, recruitment, and data collection are published elsewhere. <sup>36-38</sup>

## Baseline data collection

Information on the participants was obtained during enrolment from questionnaires that covered sociodemographic characteristics, lifestyle factors,

personal and family history of diseases, and, for women, menstrual and reproductive history. Anthropometric measurements, such as height and weight, were performed in all centres at baseline using standard procedures; except in France, the UK, and Norway, where self-reported data were collected. Updated data on weight during follow-up were obtained for a subsample of participants involved in the European Prospective Investigation into Cancer and Nutrition-Physical Activity, Nutrition, Alcohol, Cessation of Smoking, Eating Out of Home and Obesity (EPIC-PANACEA) study.<sup>39</sup>

## Dietary intake assessment

To assess the usual dietary intakes of participants, we used country specific and validated dietary questionnaires at recruitment. Depending on the study centres, these questionnaires were either selfadministered or interviewer administered semiquantitative food frequency questionnaires, with an estimation of individual average portions or with the same standard portion assigned to all participants, or diet history questionnaires, some combining a food frequency questionnaire and seven day dietary records.<sup>38</sup> The EPIC food composition database comprises more than 10000 food and beverage items reflecting the types of food consumed in each country. 40 A subset of the EPIC cohort (random samples of 5-12% of participants from each EPIC centre) also completed one computer assisted 24 hour dietary recall (EPIC-SOFT computer program), as part of a calibration study.41

# FSAm-NPS dietary index computation

The FSAm-NPS is a modified version of the original nutrient profiling system (FSA-NPS), with slight adaptations to the allocation of points for specific foods (beverages, cheese, and added fats) recommended by the French High Council for Public Health to ensure a proper discrimination of the nutritional quality of products within these groups and a high consistency of the FSAm-NPS score with nutritional recommendations. <sup>14</sup> Details on how the FSAm-NPS score is calculated are published elsewhere <sup>8</sup> <sup>12</sup> <sup>14</sup> <sup>35</sup> (also see supplementary methods).

For each food or beverage in the EPIC food composition database we calculated the FSAm-NPS score (food level score) based on its composition for each 100 g of content: we allocated A points (ie, nutrients that should be consumed in limited amounts) for total sugars (g), saturated fatty acids (g), sodium (mg), and energy (kJ) and C points (ie, nutrients or components that should be promoted) for dietary fibre (g) and protein (g) and for fruit, vegetables, legumes, and nuts (%). The percentage content of fruit, vegetables, legumes, and nuts was derived using standard recipes. A points (range 0-10 for each of the four items) and C points (range 0-5 for each of the three items) are allocated following specific grids for each item (see supplementary methods) and summed. To obtain the FSAm-NPS score the

sum of C points is then subtracted from the sum of A points (see supplementary methods). The FSAm-NPS score for each food or beverage is based on a unique discrete continuous scale ranging theoretically from –15 points (highest nutritional quality) to 40 points (lowest nutritional quality). Cut-offs are then applied to the FSAm-NPS score to derive the Nutri-Score. The supplementary methods provide examples of the FSAm-NPS score calculation, Nutri-Score cut-offs, and food products classified according to Nutri-Score.

In a second step, we calculated a FSAm-NPS dietary index to characterise the nutritional quality of an individual's diet. The FSAm-NPS dietary index (individual level score) was obtained as the sum of FSAm-NPS score for each food or beverage consumed, multiplied by the amount of energy provided by this product (energy content per 100 g multiplied by the estimated daily intake assessed using the baseline dietary questionnaires), divided by the total amount of energy intake (fig 1).<sup>42</sup> A higher FSAm-NPS dietary index score reflects an overall lower nutritional quality of foods consumed.

### Follow-up for vital status

We obtained data on vital status and cause of death through linkage to mortality registries combined with data collected during follow-up of the cohort. The end of follow-up or closure dates of the study period varied between 2012 and 2015 depending on the country. The cause of death was coded using ICD-10 (international classification of diseases, 10th revision).<sup>43</sup> In this study, in addition to mortality from all causes we considered mortality due to specific causes: cancer (C00-D48), diseases of the circulatory system (I00-199), diseases of the respiratory system (J00-J99), and diseases of the digestive system (K00-K93), and, as a negative control, mortality due to external causes (injury, poisoning, and other consequences of external causes: S00-T98, and external causes of morbidity and mortality: V01-Y98). Mortality from all non-external causes (main exposure) was defined as mortality from all causes except external causes of death.

## Statistical analyses

Of the 521324 participants, we excluded those with missing lifestyle or dietary information (n=6902), along with those with an extreme ratio of energy intake to energy requirement (highest and lowest centiles, n=10241), participants with no follow-up (n=2516), and those with missing date of death (n=71). A total of 54951 deaths were recorded during follow-up, 1839 of which were due to external causes (see flowchart in supplementary figure 1).

We calculated age standardised absolute rates as the number of cases per 10 000 persons over 10 years in the highest and lowest fifths of the FSAm-NPS dietary index.

We considered the FSAm-NPS dietary index as a continuous variable (increment of 1 standard deviation—ie, 2.1 points of score) and as sex specific fifths. Tests for linear trends were performed assigning

FSAm-NPS dietary index =  $\frac{\sum_{i=1}^{n} (FS_{i}E_{i})}{\sum_{i=1}^{n} E_{i}}$ 

FS<sub>i</sub> = score of food or beverage i E<sub>i</sub> = energy intake from food or beverage i n = number of foods or beverages consumed

Fig 1 | Equation to calculate the Food Standards Agency nutrient profiling system (FSAm-NPS) dietary index

the median for each fifth of FSAm-NPS dietary index. Restricted cubic spline modelling was used to explore non-linear associations. Cox proportional hazards regression models were computed to analyse the associations between the FSAm-NPS dietary index score and all cause and cause specific mortality. Examination of the Schoenfeld residuals confirmed that the assumptions of proportionality were satisfied (see supplementary figure 2). Participants contributed person time to the model until date of death, date of emigration or loss to follow-up, or end of followup, whichever occurred first. For analyses of cause specific mortality, participants who died from another cause than the one under study were included and censored at the date of the competing death event. Similarly, for analyses on mortality from all nonexternal causes, we included participants who died from external causes in the model and censored them at date of death. Competing risks were also tested using Fine and Gray models.44 Hazard ratios and corresponding 95% confidence intervals were derived from multivariable Cox regression models using age as the underlying time variable. Models were stratified (using Cox model stratums) by age at recruitment (one year intervals) and study centre,<sup>36</sup> to take into account a possible heterogeneity between study centres (and therefore countries). The main model accounted for all major potential confounders available through an adjustment for several covariates: sociodemographic characteristics (sex. educational level—longer education, including university degree, technical or professional school, secondary school, primary school), lifestyle (combined total physical activity—sex specific categories: active, moderately active, moderately inactive, inactive), smoking status and intensity (current, 1-15 cigarettes daily, 16-25 cigarettes daily, ≥26 cigarettes daily, and pipe, cigar, or occasional; current or former, missing; former, quit for ≤10 years, quit for 11-19 years, quit for ≥20 years; non-smoker), alcohol and energy intakes at baseline, anthropometric characteristics (body mass index (BMI), height), and prevalent disease (history of cancer, cardiovascular diseases, and diabetes).

Missing data on covariates (physical activity: n=34400 (6.9%); smoking status and intensity: n=8527 (1.7%); educational level: n=18383 (3.7%); history of cardiovascular diseases: n=78400 (15.6%), history of diabetes: n=39892 (7.9%); BMI: n=91412 (18.2%); height: n=90258 (18%)) were handled using multiple imputation by chained equations

(MICE method<sup>45</sup>) by fully conditional specification (10 imputed datasets). We also conducted a complete case approach (ie, excluding participants with missing data on covariates).

We considered BMI as a confounding factor in the analyses and therefore it was adjusted for in the multivariable models. As BMI could also be considered as an intermediate mediating factor, however, we performed sensitivity analyses without adjustment for BMI, and analyses of mediation through variation in BMI were also implemented using a method proposed previously. 46

To test the robustness of the associations, we carried out several sensitivity analyses: we removed energy intake from the models (assessing a potential collider bias), we included coffee and soft drink intakes in the models (assessing if these two dietary factors recently found to be strongly associated with mortality in EPIC<sup>47 48</sup> would entirely explain the associations), we excluded from the analyses participants with a history of cancer, cardiovascular diseases, and diabetes (assessing a potential bias from modified dietary behaviours after these major health events, such as indications to follow a healthier diet), and we excluded from the analyses those participants who died during the first five years of follow-up (allowing a longer delay between baseline dietary assessment and mortality event). To assess the potential for residual confounding we also carried out subgroup analyses according to major potential confounders (sex, BMI, physical activity, educational level, smoking status, alcohol intake, energy intake). Potential residual confounding from unmeasured confounders was assessed using E values.49 50

All tests were two sided and we considered P<0.05 to be statistically significant. SAS version 9.4 (SAS Institute) and R version 3.6.2 were used for the analyses.

### Patient and public involvement

The research question developed in this article corresponds to concerns of the participants involved in the EPIC cohort, and of the public in general. The results of the present study will be disseminated through institutional websites and the media.

# Results

A total of 501594 adults (70.8% women, median age 51.6 years) were included in this study (see supplementary figure 1). After a median follow-up of 17.2 years (8162730 person years), 54951 deaths occurred, 23143 of which were from cancer, 13246 from diseases of the circulatory system, 2857 from diseases of the respiratory system, 1561 from diseases of the digestive system, and 1839 from external causes.

Table 1 shows the baseline characteristics of the participants overall and according to sex specific fifths of the FSAm-NPS dietary index. This index reflects the overall nutritional quality of an individual's diet based on the intrinsic nutritional quality of each food consumed, regardless of cultural context. Owing

Table 1 | Baseline characteristics of study participants from European Prospective Investigation into Cancer and Nutrition (EPIC) cohort overall and by sex specific fifths of Food Standards Agency nutrient profiling system (FSAm-NPS) dietary index score. Values are numbers (percentages) unless stated otherwise

		Fifths of FSAm-NPS dietary index score†						
Characteristics	All (n=501 594)*	First (highest nutritional quality) (n=100318)‡	Second (n=100319)	Third (n=100 319)	Fourth (n=100 319)	Fifth (lowest nutritional quality (n=100319)		
Median (interquartile range)			, ,	, ,				
FSAm-NPS dietary index score	5.95 (4.53-7.39)	3.29 (2.52-3.79)	4.84 (4.53-5.13)	5.95 (5.67-6.22)	7.07 (6.78-7.39)	8.66 (8.14-9.44)		
Men	146 329 (29.2)	29 265 (20.0)	29 266 (20.0)	29 266 (20.0)	29 266 (20.0)	29 266 (20.0)		
Women	355 265 (70.8)	71 053 (20.0)	71 053 (20.0)	71 053 (20.0)	71 053 (20.0)	71 053 (20.0)		
Median (interquartile range) age at recruitment (years)	51.6 (45.3-58.4)	52.1 (45.4-59.2)	51.5 (44.9-58.3)	51.5 (44.8-58.0)	51.6 (45.4-58.3)	51.5 (45.7-58.4)		
Country:								
France	72 980 (14.5)	2558 (3.51)	6812 (9.33)	13 135 (18.0)	21795 (29.9)	28 680 (39.3)		
Italy	45 700 (9.11)	9734 (21.3)	13 945 (30.5)	11 143 (24.4)	7295 (16.0)	3583 (7.84)		
Spain	40 619 (8.10)	21 587 (53.1)	8708 (21.4)	5214 (12.8)	3098 (7.63)	2012 (4.95)		
UK	80 441 (16.0)	18 8 26 (23.4)	13 661 (17.0)	13 936 (17.3)	14652 (18.2)	19 366 (24.1)		
Netherlands	38 195 (7.61)	3736 (9.78)	7725 (20.2)	10 108 (26.5)	9999 (26.2)	6627 (17.3)		
Greece	26 651 (5.31)	10 900 (40.9)	10 032 (37.6)	4263 (16.0)	1206 (4.53)	250 (0.94)		
Germany	52010 (10.4)	4729 (9.09)	7718 (14.8)	11 107 (21.4)	13735 (26.4)	14721 (28.3)		
Sweden	52741 (10.5)	8495 (16.1)	10 311 (19.5)	11 043 (20.9)	10 982 (20.8)	11 910 (22.6)		
Denmark	55 818 (11.1)	8503 (15.2)	10 461 (18.7)	12 297 (22.0)	13 171 (23.6)	11 386 (20.4)		
Norway	36 439 (7.26)	11 250 (30.9)	10 946 (30.0)	8073 (22.1)	4386 (12.0)	1784 (4.90)		
Median (interquartile range) body mass index§	25.3 (22.8-28.2)	26.0 (23.3-29.1)	25.5 (23.1-28.5)	25.2 (22.8-28.0)	24.8 (22.5-27.6)	24.7 (22.2-27.5)		
Median (interquartile range) neight (cm)¶ Educational level**:	166 (160-173)	164 (158-170)	165 (159-172)	166 (160-173)	167 (161-174)	167 (161-174)		
None or primary school	149 580 (31.0)	40 566 (27.1)	33 650 (22.5)	27 778 (18.6)	23 976 (16.0)	23 610 (15.8)		
Technical, professional, or secondary school	214 154 (44.3)	35 580 (16.6)	41742 (19.5)	44 933 (21.0)	46 072 (21.5)	45 827 (21.4)		
Longer education (including university degree)	119 477 (24.7)	20 301 (17.0)	21 935 (18.4)	24 473 (20.5)	26 653 (22.3)	26 115 (21.9)		
Physical activity††:								
Inactive	76 414 (15.2)	12 349 (16.2)	13864 (18.1)	15 532 (20.3)	16 969 (22.2)	17 700 (23.2)		
Moderately inactive	159 880 (31.9)	26 205 (16.4)	27 882 (17.4)	30765 (19.2)	35 200 (22.0)	39 828 (24.9)		
Moderately active	185 968 (37.1)	44 635 (24.0)	40 824 (22.0)	36 879 (19.8)	33 180 (17.8)	30 450 (16.4)		
Active	44 932 (8.96)	9819 (21.8)	9429 (21.0)	9178 (20.4)	8541 (19.0)	7965 (17.7)		
Smoking status‡‡:								
Non-smoker	244 929 (48.8)	51 476 (21.0)	48 552 (19.8)	47 992 (19.6)	48 625 (19.8)	48 284 (19.7)		
Former smoker	134 382 (26.8)	27 306 (20.3)	27 400 (20.4)	27 459 (20.4)	26 815 (19.9)	25 402 (18.9)		
Current smoker	111 938 (22.3)	19756 (17.6)	22 405 (20.0)	22851 (20.4)	22817 (20.4)	24 109 (21.5)		
Median (interquartile range) alcohol intake (g/d)	5.29 (0.93-14.8)	2.86 (0.35-11.8)	4.56 (0.82-13.2)	5.56 (1.08-15.0)	6.64 (1.46-16.4)	6.81 (1.49-17.0)		
Median (interquartile range) energy (kcal/d)	1992 (1628-2430)	1745 (1432-2144)	1899 (1568-2310)	1984 (1645-2393)	2092 (1736-2502)	2253 (1863-2703)		
Median (interquartile range) total dietary fibre (g/d)	21.8 (17.4-27.0)	24.2 (19.3-30.4)	22.4 (18.1-27.5)	21.7 (17.4-26.6)	21.1 (16.9-25.9)	19.9 (15.7-24.5)		
Median (interquartile range) vegetables (g/d)	175.4 (110.0-276.3)	218.6 (133.6-339.7)	183.2 (115.3-292.1)	166.2 (107.0-260.4)	160.6 (104.0-249.3)	157.2 (98.4-243.5)		
Median (interquartile range) ruit, nuts, and seeds (g/d)	200.5 (111.7-321.6)	287.1 (173.5-434.6)	234.1 (132.1-354.5)	194.9 (111.6-308.1)	171.9 (98.8-273.2)	143.6 (80.1-234.0)		
Median (interquartile range) dairy products (g/d)	278.1 (161.3-445.9)	269.8 (146.5-448.7)	284.5 (164.5-463.6)	294.9 (173.6-465.4)	284.1 (168.2-445.5)	258.3 (153.0-400.0		
Median (interquartile range) ish and shellfish (g/d)	28.0 (13.8-49.7)	32.9 (15.1-63.6)	28.6 (14.4-53.0)	27.3 (13.6-48.9)	26.5 (13.2-44.8)	25.5 (12.8-42.3)		
Median (interquartile range) ed meat (g/d)	34.5 (16.1-62.7)	26.1 (10.1-49.9)	34.2 (16.6-60.3)	37.3 (18.0-65.9)	40.1 (19.0-69.0)	36.6 (17.1-65.9)		
Median (interquartile range) processed meat (g/d)	24.3 (10.6-43.9)	13.0 (3.22-27.4)	19.9 (7.74-36.5)	25.6 (12.5-44)	30.5 (16.1-50.9)	35.6 (18.5-59.9)		
History of cancer History of cardiovascular	24 155 (4.82) 97 370 (19.4)	4359 (18.0) 22 211 (22.8)	4456 (18.4) 19915 (20.4)	4708 (19.5) 18 997 (19.5)	5204 (21.5) 18 645 (19.2)	5428 (22.5) 17 602 (18.1)		
diseases§§ History of diabetes¶¶	13 311 (2.65)	5258 (39.5)	2853 (21.4)	2017 (15.1)	1692 (12.7)	1491 (11.2)		
				/11/115 11				

<sup>\*</sup>Column percentages.

<sup>†</sup>Cut-offs for sex specific fifths of FSAm-NPS dietary index were 4.14, 5.35, 6.43, and 7.68 for women and 4.32, 5.55, 6.63, and 7.88 for men. A higher score indicates a lower nutritional quality of foods consumed.

<sup>‡</sup>Row percentages.

<sup>§</sup>Missing for 91 412 (18.2%).

<sup>¶</sup>Missing for 90 258 (18.0%).

<sup>\*\*</sup>Missing for 18 383 (3.7%).

ttMissing for 34 400 (6.9%). ttMissing for 10 345 (2.1%).

<sup>§§</sup>Missing for 78 400 (15.6%).

<sup>¶¶</sup>Missing for 39 892 (7.9%).

to the diverse dietary patterns of the 10 countries participating in the EPIC cohort, FSAm-NPS dietary index scores were lower, indicating diets of overall higher nutritional quality in Spain (median 4.06), Greece (4.49), Norway (4.92), and Italy (5.34), and higher in the UK (6.01), Sweden (6.19), the Netherlands (6.22), Denmark (6.25), Germany (6.73), and France (7.25).

Participants with higher FSAm-NPS dietary index scores were more likely to have less healthy dietary intakes (lower intakes of dietary fibres, fruit and vegetables, and fish and higher intakes of red and processed meat) and higher energy intakes. Nonetheless, a broad range of energy intakes was observed within each fifth. Participants with higher FSAm-NPS dietary index scores were also more likely to smoke, to be less physically active, and to have a higher alcohol intake and higher level of education, and a history of cancer. In contrast, a higher proportion of existing cardiovascular diseases or diabetes and a higher BMI was observed in participants with lower FSAm-NPS dietary index scores, which might reflect a change in diet required for management of disease in these participants.

Table 2 shows the associations between the FSAm-NPS dietary index and mortality. A higher score (lower nutritional quality diet) was associated with higher mortality overall (highest fifth versus lowest fifth: hazard ratio 1.06, 95% confidence interval 1.03 to 1.09, P<0.001 for trend, P<0.001 for non-trend). Mortality from external causes was not associated with the FSAm-NPS dietary index score (0.99, 0.84 to 1.16, P=0.9 for trend, P=0.9 for non-trend). Mortality from all non-external causes (all cause mortality) was positively associated with the FSAm-NPS dietary index (n=53112 cases, 1.07, 1.03 to 1.10, P<0.001 for trend, P<0.001 for non-trend). Corresponding age standardised absolute rates for all cause mortality per 10000 persons over 10 years were 760 (men=1237; women=563) in those with a high FSAm-NPS dietary index (low nutritional quality diet) and 661 (men=1008; women=518) in those with low scores (high nutritional quality diet).

Overall, results were consistent across all countries (see supplementary figure 3), with only borderline statistically significant associations (restricted sample size). Results for Norway might be related to the distribution of the FSAm-NPS dietary index scores in this country, with overall low scores and a resulting small contrast between individuals with higher and lower scores (median 4.92, interquartile range 3.87-5.98).

Cause specific analyses showed that a higher FSAm-NPS dietary index score (lower nutritional quality diet) was associated with higher mortality from cancer (highest fifth versus lowest fifth: hazard ratio 1.08, 1.03 to 1.13, P<0.001 for trend, P=0.003 for non-trend) and diseases of the circulatory (1.04, 0.98 to 1.11, P=0.06 for trend, P=0.02 for non-trend), respiratory (1.39, 1.22 to 1.59, P<0.001 for trend, P<0.001 for non-trend), and digestive systems (1.22, 1.02 to 1.45, P=0.03 for trend, P=0.19 for non-trend).

Some evidence of non-linearity was observed for all cause mortality and mortality from cancer and diseases of the circulatory system. Such non-linearity was mainly observed for low FSAm-NPS dietary index scores, whereas for higher scores the association had a linear shape (see supplementary figure 4). Similar results were observed across all sensitivity analyses (see supplementary table 1).

Overall, subgroup analyses showed the robustness of the results across categories of major potential confounders (fig 2): associations with all cause mortality were consistent across stratums for men and women, non-smokers and smokers, and according to energy or alcohol intakes and education levels (although strengthened in highly educated participants); associations were stronger in non-obese participants and in those who were less physically active.

Mediation analyses suggested a limited mediation effect from variation in BMI during follow-up in the association between FSAm-NPS dietary index score and mortality (see supplementary table 2). Removing BMI from the models did not change the results of the main models (see supplementary table 1).

Finally, Evalues suggested that residual confounding due to potential unmeasured confounding factors is likely to be moderate (see supplementary table 3).

#### **Discussion**

This study was conducted in a large population from 10 European countries participating in the EPIC cohort to assess the relevance of the FSAm-NPS dietary index score (high values representing low nutritional quality of food products) to characterise healthier food choices in a European context. We found that a higher consumption of food products with higher FSAm-NPS scores (ie, higher FSAm-NPS dietary index scores at the individual level) was positively associated with mortality from all causes and from cancer and diseases of the circulatory, respiratory, and digestive systems.

## Comparison with other studies

This work builds on previous analyses conducted on cancer incidence in the EPIC cohort<sup>35</sup> to investigate the association between the FSAm-NPS dietary index and health outcomes in a large European population, and it complements the analyses conducted in the French SU.VI.MAX and NutriNet-Santé cohorts.<sup>27</sup> <sup>29-34</sup> These studies consistently reported poorer health outcomes (weight gain,<sup>29</sup> metabolic syndrome,<sup>30</sup> cancer,<sup>33-35</sup> cardiovascular diseases,<sup>31</sup> <sup>32</sup> asthma symptoms<sup>27</sup>) associated with higher FSAm-NPS dietary index scores.

Previous studies in the UK have also investigated the association between the FSA-NPS score (the original score before modifications were made and it was renamed FSAm-NPS) and mortality, applying to the FSA-NPS a cut-off to categorise food products as healthier or as less healthy (Ofcom threshold used for advertising regulation<sup>9</sup>). The results of these studies were consistent with ours, showing a lower all cause and cancer related mortality associated with intake of

Table 2 | Associations between fifths of Food Standards Agency nutrient profiling system (FSAm-NPS) dietary index score and all cause and cause specific mortality, from multivariable Cox proportional hazards regression models, in participants of European Prospective Investigation into Cancer and Nutrition (EPIC) cohort, 1992-2015. Values are hazard ratios (95% confidence intervals) unless stated otherwise

Mortality	Continuous	P value	Fifths*					P for	
	(per 1 SD increment)		First (highest nutritional quality)	Second	Third	Fourth	Fifth (lowest nutritional quality)	P for trend	non- trend
All causes									
No/person years	54951/8162730		10887/1605206	9934/1585846	10 275/1 626 056	11 098/1 662 098	12757/1683523		
Sex adjusted model†	1.04 (1.03 to 1.05)	<0.001		0.97 (0.94 to 1.00)	0.98 (0.95 to 1.01)	1.01 (0.98 to 1.04)	1.10 (1.07 to 1.14)	<0.001	<0.001
Main model‡	1.02 (1.01 to 1.03)	<0.001	1.00 (ref)	0.98 (0.96 to 1.01)	0.99 (0.96 to 1.02)	1.01 (0.98 to 1.04)	1.06 (1.03 to 1.09)	<0.001	<0.001
Cause specific									
Non-external:									
No/person years	53 112/8 162 730		10515/1605206	9605/1585846	9922/1626056	10728/1662098	12342/1683523		
Sex adjusted model	1.04 (1.03 to 1.05)	<0.001	1.00 (ref)	0.97 (0.94 to 1.00)	0.98 (0.95 to 1.01)	1.01 (0.98 to 1.04)	1.10 (1.07 to 1.14)	<0.001	<0.001
Main model	1.03 (1.02 to 1.04)	<0.001	1.00 (ref)	0.99 (0.96 to 1.02)	0.99 (0.96 to 1.02)	1.01 (0.98 to 1.04)	1.07 (1.03 to 1.1)	<0.001	<0.001
External:									
No/person years	1839/7783132		372/1568430	329/1538426	353/1556361	370/1562787	415/1557129		
Sex adjusted model	1.03 (0.98 to 1.09)	0.23		0.95 (0.81 to 1.10)	1.00 (0.86 to 1.17)	1.00 (0.86 to 1.17)	1.08 (0.92 to 1.27)	0.21	0.54
Main model	1.00 (0.95 to 1.05)	0.93	1.00 (ref)	0.94 (0.81 to 1.1)	0.98 (0.84 to 1.15)	0.96 (0.82 to 1.13)	0.99 (0.84 to 1.16)	0.98	0.93
Cancer:									
No/person years	23 143/7 783 132		4550/1568430	4288/1538426	4482/1556361	4700/1562787	5123/1 557 129		
Sex adjusted model	1.06 (1.04 to 1.07)	<0.001	1.00 (ref)	1.00 (0.96 to 1.04)	1.04 (0.99 to 1.08)	1.07 (1.03 to 1.12)	1.16 (1.11 to 1.21)	<0.001	<0.001
Main model	1.03 (1.01 to 1.04)	<0.001	1.00 (ref)	0.99 (0.95 to 1.04)	1.02 (0.98 to 1.07)	1.03 (0.99 to 1.08)	1.08 (1.03 to 1.13)	<0.001	0.003
Circulatory diseases:									
No/person years	13 246/7 783 132		2973/1568430	2432/1538426	2377/1556361	2526/1562787	2938/1557129		
Sex adjusted model	1.02 (1.00 to 1.04)	0.04	1.00 (ref)	0.91 (0.86 to 0.96)	0.92 (0.87 to 0.97)	0.95 (0.90 to 1.01)	1.03 (0.97 to 1.09)	0.11	<0.001
Main model	1.02 (1.00 to 1.04)	0.03	1.00 (ref)	0.96 (0.91 to 1.01)	0.96 (0.91 to 1.02)	1.00 (0.94 to 1.06)	1.04 (0.98 to 1.11)	0.06	0.02
Respiratory diseases:									
No/person years	2857/7783132		508/1568430	501/1538426	507/1556361	591/1562787	750/1557129		
Sex adjusted model	1.16 (1.12 to 1.21)	<0.001	1.00 (ref)	1.12 (0.98 to 1.27)	1.15 (1.01 to 1.31)	1.30 (1.14 to 1.47)	1.56 (1.37 to 1.76)	<0.001	<0.001
Main model	1.11 (1.06 to 1.15)	<0.001	1.00 (ref)	1.15 (1.01 to 1.31)	1.16 (1.01 to 1.32)	1.27 (1.11 to 1.45)	1.39 (1.22 to 1.59)	<0.001	<0.001
Digestive diseases:									
No/person years	1561/7783132		294/1568430	286/1538426	282/1556361	326/1562787	373/1557129		
Sex adjusted model	1.09 (1.03 to 1.15)	0.002	1.00 (ref)	1.06 (0.89 to 1.25)	1.03 (0.87 to 1.22)	1.15 (0.97 to 1.36)	1.25 (1.06 to 1.48)	0.005	0.05
Main model	1.08 (1.02 to 1.14)	0.01	1.00 (ref)	1.08 (0.91 to 1.28)	1.05 (0.88 to 1.25)	1.15 (0.97 to 1.37)	1.22 (1.02 to 1.45)	0.03	0.19

<sup>\*</sup>Cut-offs for sex specific fifths of the FSAm-NPS dietary index were 4.14, 5.35, 6.43, and 7.68 for women and 4.32, 5.55, 6.63, and 7.88 for men. A higher score indicates a lower nutritional quality of foods consumed.

a wide variety of healthier food items in the Whitehall II cohort<sup>51</sup> and a higher all cause mortality associated with a higher consumption of less healthy food items in EPIC-Norfolk (highest versus lowest fifth: hazard ratio 1.11, 95% confidence interval 1.02 to 1.20).<sup>52</sup> In both studies no association was observed with mortality from cardiovascular diseases, whereas in our study we found a borderline statistically significant association. This might be related to the larger sample size in our study or to a better ranking of participants using the continuous FSAm-NPS, which allows for a

more refined discrimination of the nutritional quality of food products likely resulting in a better ranking of participants according to the overall nutritional quality of their diets.

The approach used in our study, in which a dietary index at the individual level is derived from the nutrient profile of the foods consumed and is studied in relation to health outcomes, was also implemented with different nutrient profiling systems in two other studies. In the Nurses' Health Study and the Health Professionals Follow-up Study, the Overall Nutritional

<sup>&#</sup>x27;Sex adjusted model was stratified for age (one year interval) and study centre and adjusted for age (time scale) and sex.

<sup>‡</sup>The main model was stratified for age (one year interval) and study centre and adjusted for age (time scale) sex, body mass index, height, educational level (longer education, including university degree, technical or professional school, secondary school, primary school, missing), combined total physical activity (sex specific categories: active, moderately active, inactive, missing), smoking status and intensity of smoking (current, 1-15 cigarettes daily; 16-25 cigarettes daily; ≥26 cigarettes daily; pipe, cigar, occasional; current or former, missing; former, quit 11-19 years, quit ≥20 years, quit ≤10 years; non-smoker; missing), baseline alcohol intake, baseline energy intake, and history of cancer (yes, no), cardiovascular diseases (yes, no, missing) and diabetes (yes, no, missing).

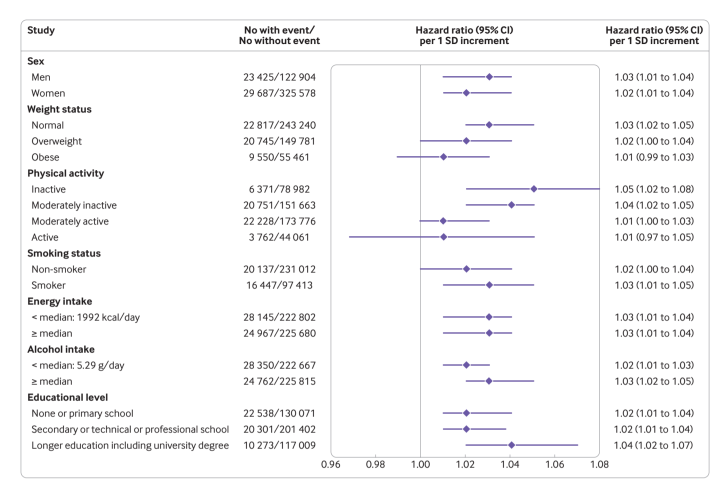


Fig 2 | Associations between Food Standards Agency nutrient profiling system (FSAm-NPS) dietary index score and all cause mortality, subgroup analyses from multivariable Cox proportional hazards regression models in the European Prospective Investigation into Cancer and Nutrition (EPIC) cohort, 1992-2015. A higher score indicates a lower nutritional quality of consumed foods. The main model was stratified for age (one year interval) and study centre and adjusted for sex, body mass index, height, educational level (longer education, including university degree, technical or professional school, secondary school, primary school, missing), combined total physical activity (sex specific categories: active, moderately active, moderately inactive, inactive, missing), smoking status and intensity of smoking (current, 1-15 cigarettes daily, 16-25 cigarettes daily, ≥26 cigarettes daily, pipe, cigar, or occasional; current or former, missing; former, quit ≤10 years, quit 11-19 years, quit ≥20 years; non-smoker; missing), baseline alcohol intake, baseline energy intake, and history of cancer (yes, no), cardiovascular diseases (yes, no, missing), and diabetes (yes, no, missing). P for interaction, obtained for each subgroup analysis from the likelihood ratio test of models with and without the interaction term, were: sex, P=0.04; weight status, P=0.22; physical activity, P=0.23; smoking status, P<0.001; energy intake, P=0.05; alcohol intake, P=0.07; educational level. P=0.27

Quality Index translated into the ONQI-f score at the individual level was associated with lower mortality.<sup>53</sup> In the Rotterdam Study, the Nutrient-Rich Food 9.3 score at the individual level was inversely associated with all cause mortality but not with mortality from cardiovascular diseases.54 These scores nonetheless differ by the number and types of nutritional items considered. The FSAm-NPS was designed to be easily computable by industrial and public stakeholders in a transparent manner to serve as a basis for tools of public health nutritional policies (such as the Nutri-Score label). Hence the FSAm-NPS consists of a unique scale applicable to all food products (raw or manufactured) and to all countries (as was done in the present study) and intentionally focuses on seven items only (energy, saturated fatty acids, sodium, sugars, dietary fibre, and protein, and fruit, vegetables, legumes, and nuts). These items are generally found in the nutritional

information on food labels and were selected based on an association with non-communicable diseases or because they reflect the nutritional value of foods, in line with dietary guidelines. All In contrast, the Overall Nutritional Quality Index is based on 30 items and the Nutrient-Rich Food 9.3 is based on 12 items, both including not only macronutrients, vitamins, and minerals but also polyphenols (Overall Nutritional Quality Index) and reference values of intake (Nutrient-Rich Food 9.3) that might differ across countries.

The FSAm-NPS is also consistent with recent reports from the Global Burden of Disease Study and the EAT-Lancet Commission, both of which estimated that about 11 million deaths worldwide could be prevented with healthier diets, including, notably, less sodium, sugars, and saturated fats and more dietary fibre and whole grains, fruit, vegetables, legumes, and nuts. <sup>155</sup> Our results showed a strong association of the FSAm-

NPS dietary index score with mortality from respiratory diseases. Beyond the well established impact of nutrition on cancer and cardiometabolic risks,1 mounting evidence also supports a substantial impact of nutrition on respiratory health through several pathways involving oxidative stress and inflammation. epigenetics, and the gut microbiome. Notably, dietary fibres (involved in anti-inflammatory responses) and fruit and vegetables (sources of antioxidants), as part of a healthy diet, have been suggested to play a beneficial role in respiratory health, whereas components such as saturated fats and red or processed meat (involved in pro- inflammatory responses), or, more generally, a Western diet, would have detrimental effects. 56-61 The FSAm-NPS dietary index score has also been associated with asthma symptoms in the NutriNet-Santé cohort study.<sup>27</sup> Similar strong associations with mortality from respiratory diseases (compared with mortality from other causes) have been observed in previous studies of saturated fatty acid<sup>62</sup> or vegetables and red meat intakes.<sup>63</sup> These results are consistent with the items in the FSAm-NPS dietary index.

# Non-linearity and associations between FSAm-NPS dietary index and BMI

In our study, evidence of non-linearity was observed for associations with all cause mortality and mortality from cancer and diseases of the circulatory system at low values of FSAm-NPS dietary index scores. Such values reflect healthy food choices that might have been adopted by individuals with a greater risk of disease (and thus high underlying risks of mortality). Hence, individuals consuming diets of the highest nutritional quality (lowest FSAm-NPS dietary index scores) would be the ones with higher mortality rates, thus blurring the association overall. Evidence of this can be seen in table 1 where participants in the lowest fifth of the FSAm-NPS dietary index have a higher baseline BMI (cross sectional analyses) and were more likely to have prevalent cardiovascular disease or prevalent diabetes at baseline (probably partly related to their past diet). This might also help to explain why weaker (although statistically significant) associations were observed for mortality from circulatory diseases or why we observed stronger results in non-obese participants at baseline, because mechanisms leading to premature death have to some extent already played out for obese individuals. Finally, our analyses showed little mediation effect of variation in BMI in the association between the FSAm-NPS dietary index score and mortality. This suggests that the overall nutritional quality of a diet might have an impact on mortality beyond weight gain.

# Associations between FSAm-NPS dietary index and educational level

The positive cross sectional association between the FSAm-NPS dietary index score and educational level might seem counterintuitive but has also been observed in studies conducted in the independent SU.VI.MAX and NutriNet-Santé cohorts. 42 64 Several hypotheses can explain this finding. In EPIC, countries with a higher proportion of participants with lower mean educational level (eg. Greece, Spain, Italy) also had the lowest FSAm-NPS dietary index scores (reflecting trends towards a Mediterranean or healthier diet in countries of southern Europe<sup>65</sup> 66). At least in these countries, people with lower education might be more likely to consume traditional diets that could include food of better nutritional quality. This trend could also be related to an age or generation effect. Younger people tend to have a higher educational level but also to have a higher consumption of "junk food," leading to higher FSAm-NPS dietary index scores (diets of lower nutritional quality). Associations were statistically significant for all categories of educational level but slightly strengthened in highly educated participants. The latter finding might be related to more contrasted FSAm-NPS dietary index scores in the category of participants with longer education. Another hypothesis is that participants with longer education might be less exposed to risk factors linked to occupation or environment than those of a lower socioeconomic position.

### Strengths and limitations of this study

Strengths of our study pertain to its prospective design and long follow-up of many participants from different European countries with various phenotypes, and for whom collected data have been standardised. This provided an opportunity to study the nutritional quality of food choices in relation to mortality in a broad European context of diverse dietary patterns. Additionally, a large array of lifestyle data was available which allowed adjustment for major potential confounding factors in our main model and the performance of in-depth sensitivity analyses.

Our study also has some limitations. Firstly, owing to the observational design, potential residual confounding or unmeasured confounding (from genetic or environmental factors that could not be taken into account) cannot be ruled out. In addition, as we collected data using questionnaires, we cannot exclude misclassification bias from imprecise dietary data and covariates. Because data were collected before the studied outcome (prospective design), any misclassification resulting from measurement errors is likely to have been non-differential-that is, independent of death status. Nonetheless, this might have resulted in biased estimates of effect (underestimation or overestimation). 67 68 In particular, the tools used to estimate an individual's usual diet are subject to imprecision and inaccuracy (such as misreporting bias, inherent in assessments of usual dietary intakes) and therefore could have resulted in some misclassification of the nutritional quality of foods consumed by participants. Indeed, most EPIC centres used a food frequency questionnaire to assess dietary intakes, which, despite allowing for a good estimation of usual dietary intakes, still limits discrimination between the nutritional quality of individual food products (compared with, for example, repeated 24 hour dietary records, as used

in the SU.VI.MAX and NutriNet-Santé cohorts, where larger effect estimates were observed for associations between FSAm-NPS dietary index scores and health outcomes<sup>31-34</sup> 61). In the subsample from the EPIC calibration study (n=34367), good concordance was shown between the FSAm-NPS dietary index score calculated from the dietary questionnaires (mainly food frequency questionnaires) and the calibrated 24 hour dietary recall (68% of participants were classified in either the same or the adjacent fifth, only 3% were classified in extreme opposite fifths, data not tabulated). However, only one day of 24 hour recall was available for participants in the calibration study. Therefore, although the dietary questionnaires available for the whole EPIC cohort might have some limitations about the level of detail available for each food item, the questionnaires still provide a better overview of the usual diet of participants compared with the single 24 hour recall. In addition, we only assessed dietary intakes at baseline. Although changes in food consumption might have occurred during followup, it is hypothesised that the baseline estimation usually reflects general eating behaviour throughout middle age. 69 Also, EPIC participants were volunteers involved in a long term cohort study investigating the association between nutrition and health and likely had more health conscious behaviours and less unhealthy dietary behaviours than the general population. This might have resulted in weaker observed associations (hence a smaller effect size) owing to smaller differences between high and low FSAm-NPS dietary index scores. Additionally, EPIC participants were recruited from 10 countries in western Europe and so caution is warranted in extrapolating the results to other populations or ethnicities worldwide. Finally, the order of magnitude for the association between the FSAm-NPS dietary index score and mortality found in our study was relatively modest but still in line with the one traditionally observed in nutritional epidemiology, and it was similar to the one observed in the study on cancer incidence in EPIC.<sup>35</sup> From a public health perspective, the opportunity to prevent 7% of premature deaths globally through healthier food choices might nonetheless be of great interest.

Potential sources of bias in our study warrant caution in the interpretation of the findings. Despite the low hazard ratios, several things allow for some confidence in our results and are in favour of an association supporting possible causality, beyond residual confounding: the robustness of associations across sensitivity analyses, including across categories of major potential confounders; null results obtained for associations between FSAm-NPS dietary index scores and mortality from external causes, which are unrelated to diet (negative control); Evalues that suggest moderate potential unmeasured confounding<sup>49</sup> <sup>50</sup>; consistency of the results with previous studies on the association between the FSAm-NPS dietary index score and health outcomes in independent cohorts<sup>27</sup> <sup>29-34</sup>; and consistency of the results with mechanistic hypotheses, supporting biological plausibility.

### Conclusions and policy implications

In this large cohort study involving 10 European countries, a diet composed on average of more food products with higher FSAm-NPS scores, which reflected poorer nutritional profiles, was associated with higher all cause and cause specific mortality. Overall, that a higher FSAm-NPS dietary index score, obtained for varied dietary patterns in different countries representing a diverse European population, leads to higher mortality rates, suggests that the FSAm-NPS is a relevant tool to characterise more or less healthy food products, no matter the food category or the specificities of the national dietary patterns.

This study adds to the current body of evidence for the FSAm-NPS score and for Nutri-Score, a nutrition label derived from the FSAm-NPS: studies linking the FSAm-NPS dietary index to health outcomes, including one study in the EPIC cohort, 27 29-35 and studies on the perception and understanding of Nutri-Score and its actual impact on food choices. 24-26 28 70 Together, these results back up the relevance of using the FSAm-NPS to grade the nutritional quality of food products in the framework of public health nutritional measures such as the Nutri-Score label, a tool aimed at the general public and patients to help them to choose food products of a higher nutritional quality. This is important considering ongoing and future debates at the EU level about making food labelling systems uniform on the front of food product packaging.

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We thank the EPIC participants and staff for contributing to this study. This work was performed within the framework of the French Network for Nutrition And Cancer Research (https://www6.inra.fr/nacre/). When authors are identified as staff of the International Agency for Research on Cancer/World Health Organization, the authors alone are responsible for the views expressed in this article and they do not necessarily represent the decisions, policy, or views of IARC/WHO.

Contributors: MD, MT, SH, CJ, and IH conceived the study and defined the analytical strategy. MD, MT, and IH performed statistical analyses and provided preliminary interpretation of findings. MD and the writing group (IH, CJ, SH, ME, MJG, MT) drafted the manuscript. CB, CC, HAW, KO, ATJ, ALR-H, M-CB-R, FRM, YM-S, TK, VK, MMB, MBS, ATr, AK, EP, GM, CA, MSDM, RT, CS, JMAB, WMMV, YTVdS, GS, TB, MLR, AA, DP, SMC-Y, AB, PA, ES, UE, JO, BSu, NJW, NGF, PV, KKT, AKn, KP, PF, ER, and MJG acquired the data and obtained funding to continue the study. All authors critically interpreted the results, revised the manuscript, provided relevant intellectual input, and read and approved the final manuscript. MD and MT had primary responsibility for the final content and are the guarantors. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted.

Funding: This work was funded by a research grant from the French National Cancer Institute (INCa)-Cancéropôle Ile-de-France (convention No 2017-1-PL SHS-01-INSERM ADR 5-1, principal investigator: MT, co-principal investigator MD). MD was supported by a grant from the Fondation Recherche Médicale (ARF201809007046). The coordination of EPIC is financially supported by the European Commission (DG-SANCO) and the International Agency for Research

on Cancer. The national cohorts are supported by the Danish Cancer Society (Denmark); Ligue Contre le Cancer, Institut Gustave Roussy, Mutuelle Générale de l'Education Nationale, and Institut National de la Santé et de la Recherche Médicale (Inserm), (France); Deutsche Krebshilfe, Deutsches Krebsforschungszentrum, and Federal Ministry of Education and Research (Germany): the Hellenic Health Foundation (Greece); Associazione Italiana per la Ricerca sul Cancro-AIRC-Italy and National Research Council (Italy); Dutch Ministry of Public Health, Welfare, and Sports, Netherlands Cancer Registry, LK Research Funds, Dutch Prevention Funds, Dutch ZON (Zorg Onderzoek Nederland), World Cancer Research Fund, and Statistics Netherlands (the Netherlands); Health Research Fund, Instituto de Salud Carlos III, regional governments of Andalucía, Asturias, Basque Country, Murcia, and Navarra, and the Catalan Institute of Oncology (Spain); Swedish Cancer Society, Swedish Scientific Council, and county councils of Skåne and Västerbotten (Sweden); Cancer Research UK (C864/A14136 to EPIC-Norfolk; C8221/A19170 to EPIC-Oxford), Medical Research Council (MR/N003284/1 and MC-UU 12015/1 to EPIC-Norfolk: MR/M012190/1 to EPIC-Oxford: UK). NJW and NGF acknowledge funding from the Medical Research Council Epidemiology Unit (MC\_UU\_12015/1 and MC\_UU\_12015/5) and National Institute for Health Research Biomedical Research Centre, Cambridge: Nutrition, Diet, and Lifestyle Research Theme (IS-BRC-1215-20014). AK and KP are supported by the Wellcome Trust, Our Planet, and Our Health (Livestock, Environment and People) (205212/Z/16/Z). The funders had no role in the study design or in the collection, analysis, interpretation of data, writing of the report, or decision to submit the article for publication.

Competing interests: All authors have completed the ICMJE uniform disclosure form at www.icmje.org/coi\_disclosure.pdf and declare: support from the French National Cancer Institute (INCa)-Cancéropôle lle-de-France and the Fondation Recherche Médicale; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years; no other relationships or activities that could appear to have influenced the submitted work.

**Ethical approval:** This study was approved by the local ethics committees and the internal review board of the International Agency for Research on Cancer.

**Data sharing:** Information on submitting an application for access to EPIC data or biospecimens is available at https://epic.iarc.fr/access/index.php.

The lead authors (the manuscript's guarantors) affirm that the manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

Dissemination to participants and related patient and public communities: The results of this article will be disseminated as lay summaries to the public through different channels, including articles on the open web platform The Conversation (in English and French), press releases, social media accounts of our research team, International Agency for Research on Cancer newsletter, and communications through the French Nutrition and Cancer Research network (www.inra.fr/nacre).

**Provenance and peer review:** Not commissioned; externally peer reviewed.

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- Afshin A, Sur PJ, Fay KA, et al, GBD 2017 Diet Collaborators. Health effects of dietary risks in 195 countries, 1990-2017: a systematic analysis for the Global Burden of Disease Study 2017. *Lancet* 2019;393:1958-72. doi:10.1016/S0140-6736(19)30041-8
- Organisation for Economic Co-operation and Development. Promoting sustainable consumption—good practices in OECD countries. OECD, Paris, 2008. www.oecd.org/ greengrowth/40317373.pdf
- 3 Hercberg S. Report to the French Ministry of Health: Propositions pour un nouvel élan de la politique nutritionnelle française de santé publique dans le cadre de la Stratégie Nationale de Santé 1ère Partie: Mesures concernant la Prévention nutritionnelle. 2013. https://solidarites-sante.gouv.fr/IMG/pdf/rapport\_ Hercberg\_15\_11\_2013.pdf

- 4 Hawley KL, Roberto CA, Bragg MA, Liu PJ, Schwartz MB, Brownell KD. The science on front-of-package food labels. *Public Health Nutr* 2013;16:430-9. doi:10.1017/S1368980012000754
- 5 Hersey JC, Wohlgenant KC, Arsenault JE, Kosa KM, Muth MK. Effects of front-of-package and shelf nutrition labeling systems on consumers. *Nutr Rev* 2013;71:1-14. doi:10.1111/nure.12000
- 6 Hercberg S. Report to the French Ministry of Health: 'Propositions pour un nouvel élan de la politique nutritionnelle française de santé publique dans le cadre de la Stratégie Nationale de Santé 1ère Partie: Mesures concernant la Prévention nutritionnelle'. 2013. https://solidarites-sante.gouv.fr/IMG/pdf/rapport\_ Hercberg\_15\_11\_2013.pdf
- 7 Arambepola C, Scarborough P, Rayner M. Validating a nutrient profile model. *Public Health Nutr* 2008;11:371-8. doi:10.1017/ S1368980007000377
- 8 Rayner M, Scarborough P, Stockley P, et al. Nutrient profiles: Development of Final Model. FSA, 2005.
- 9 Rayner M, Scarborough P, Lobstein T. The UK Ofcom Nutrient Profiling Model: Defining 'healthy' and 'unhealthy' foods and drinks for TV advertising to children. London: Ofcom 2009. www.ndph.ox.ac.uk/ cpnp/files/about/uk-ofcom-nutrient-profile-model.pdf
- 10 Arrêté du 31 Octobre 2017 fixant la forme de présentation complémentaire à la déclaration nutritionnelle recommandée par l'Etat en application des articles L. 3232-8 et R. 3232-7 du code de la santé publique [in French]. Journal Officiel de la République Française 2017:0257.
- Julia C, Etilé F, Hercberg S. Front-of-pack Nutri-Score labelling in France: an evidence-based policy. *Lancet Public Health* 2018;3:e164. doi:10.1016/S2468-2667(18)30009-4
- Julia C, Kesse-Guyot E, Touvier M, Méjean C, Fezeu L, Hercberg S. Application of the British Food Standards Agency nutrient profiling system in a French food composition database. *Br J Nutr* 2014;112:1699-705. doi:10.1017/S0007114514002761
- 13 Anses. Evaluation de la faisabilité du calcul d'un score nutritionnel tel qu'élaboré par Rayner et al. Rapport d'appui scientifique et technique. Paris: Anses, 2015. www.anses.fr/fr/system/files/ DER2014sa0099Ra.pdf
- 14 Haut Conseil de la Santé Publique. *Opinion on information regarding* the nutritional quality of foodstuffs. 2015.
- 15 Ducrot P, Méjean C, Julia C, et al. Effectiveness of Front-Of-Pack Nutrition Labels in French Adults: Results from the NutriNet-Santé Cohort Study. PLoS One 2015;10:e0140898. doi:10.1371/journal. pone.0140898
- Ducrot P, Méjean C, Julia C, et al. Objective Understanding of Frontof-Package Nutrition Labels among Nutritionally At-Risk Individuals. Nutrients 2015;7:7106-25. doi:10.3390/nu7085325
- 17 Ducrot P, Julia C, Méjean C, et al. Impact of Different Front-of-Pack Nutrition Labels on Consumer Purchasing Intentions: A Randomized Controlled Trial. Am J Prev Med 2016;50:627-36. doi:10.1016/j. amepre.2015.10.020
- Julia C, Kesse-Guyot E, Ducrot P, et al. Performance of a five category front-of-pack labelling system - the 5-colour nutrition label - to differentiate nutritional quality of breakfast cereals in France. BMC Public Health 2015;15:179. doi:10.1186/s12889-015-1522-y
- Julia C, Ducrot P, Péneau S, et al. Discriminating nutritional quality of foods using the 5-Color nutrition label in the French food market: consistency with nutritional recommendations. *Nutr J* 2015;14:100. doi:10.1186/s12937-015-0090-4
- 20 Julia C, Blanchet O, Méjean C, et al. Impact of the front-of-pack 5-colour nutrition label (5-CNL) on the nutritional quality of purchases: an experimental study. Int J Behav Nutr Phys Act 2016;13:101. doi:10.1186/s12966-016-0416-4
- 21 Julia C, Hercberg S. Research and lobbying conflicting on the issue of a front-of-pack nutrition labelling in France. *Arch Public Health* 2016;74:51. doi:10.1186/s13690-016-0162-8
- 22 Julia C, Méjean C, Péneau S, et al. The 5-CNL Front-of-Pack Nutrition Label Appears an Effective Tool to Achieve Food Substitutions towards Healthier Diets across Dietary Profiles. PLoS One 2016;11:e0157545. doi:10.1371/journal.pone.0157545
- 23 Julia C, Péneau S, Buscail C, et al. Perception of different formats of front-of-pack nutrition labels according to sociodemographic, lifestyle and dietary factors in a French population: cross-sectional study among the NutriNet-Santé cohort participants. BMJ Open 2017;7:e016108. doi:10.1136/bmjopen-2017-016108
- 24 Julia C, Hercberg S. Development of a new front-of-pack nutrition label in France: the five-colour Nutri-Score. *Public Health Panorama* 2017;3:712-25.
- 25 Egnell M, Kesse-Guyot E, Galan P, et al. Impact of Front-of-Pack Nutrition Labels on Portion Size Selection: An Experimental Study in a French Cohort. Nutrients 2018;10:1268. doi:10.3390/nu10091268
- 26 Egnell M, Ducrot P, Touvier M, et al. Objective understanding of Nutri-Score Front-Of-Package nutrition label according to individual characteristics of subjects: Comparisons with other format labels. PLoS One 2018;13:e0202095. doi:10.1371/journal.pone.0202095

- 27 Andrianasolo RM, Julia C, Varraso R, et al. Association between an individual dietary index based on the British Food Standard Agency Nutrient Profiling System and asthma symptoms. Br J Nutr 2019;12:2:63-70. doi:10.1017/S0007114519000655
- 28 Egnell M, Talati Z, Hercberg S, Pettigrew S, Julia C. Objective Understanding of Front-of-Package Nutrition Labels: An International Comparative Experimental Study across 12 Countries. Nutrients 2018;10:1542. doi:10.3390/nu10101542
- 29 Julia C, Ducrot P, Lassale C, et al. Prospective associations between a dietary index based on the British Food Standard Agency nutrient profiling system and 13-year weight gain in the SU.VI.MAX cohort. Prev Med 2015;81:189-94. doi:10.1016/j.ypmed.2015.08.022
- 30 Julia C, Fézeu LK, Ducrot P, et al. The Nutrient Profile of Foods Consumed Using the British Food Standards Agency Nutrient Profiling System Is Associated with Metabolic Syndrome in the SU.VI.MAX Cohort. J Nutr 2015;145:2355-61. doi:10.3945/jn.115.213629
- 31 Adriouch S, Julia C, Kesse-Guyot E, et al. Prospective association between a dietary quality index based on a nutrient profiling system and cardiovascular disease risk. *Eur J Prev Cardiol* 2016;23:1669-76. doi:10.1177/2047487316640659
- 32 Adriouch S, Julia C, Kesse-Guyot E, et al. Association between a dietary quality index based on the food standard agency nutrient profiling system and cardiovascular disease risk among French adults. *Int J Cardiol* 2017;234:22-7. doi:10.1016/j. ijcard.2017.02.092
- 33 Donnenfeld M, Julia C, Kesse-Guyot E, et al. Prospective association between cancer risk and an individual dietary index based on the British Food Standards Agency Nutrient Profiling System. Br J Nutr 2015;114:1702-10. doi:10.1017/S0007114515003384
- 34 Deschasaux M, Julia C, Kesse-Guyot E, et al. Are self-reported unhealthy food choices associated with an increased risk of breast cancer? Prospective cohort study using the British Food Standards Agency nutrient profiling system. BMJ Open 2017;7:e013718. doi:10.1136/bmjopen-2016-013718
- 35 Deschasaux M, Huybrechts I, Murphy N, et al. Nutritional quality of food as represented by the FSAm-NPS nutrient profiling system underlying the Nutri-Score label and cancer risk in Europe: Results from the EPIC prospective cohort study. PLoS Med 2018;15:e1002651. doi:10.1371/journal.pmed.1002651
- 36 Ferrari P, Day NE, Boshuizen HC, et al. The evaluation of the diet/ disease relation in the EPIC study: considerations for the calibration and the disease models. *Int J Epidemiol* 2008;37:368-78. doi:10.1093/ije/dym242
- 37 Riboli E, Kaaks R. The EPIC Project: rationale and study design. European Prospective Investigation into Cancer and Nutrition. Int J Epidemiol 1997;26(Suppl 1):S6-14. doi:10.1093/ije/26.suppl 1.S6
- 38 Riboli E, Hunt KJ, Slimani N, et al. European Prospective Investigation into Cancer and Nutrition (EPIC): study populations and data collection. *Public Health Nutr* 2002;5(6B):1113-24. doi:10.1079/ PHN2002394
- 39 Vergnaud A-C, Norat T, Romaguera D, et al. Meat consumption and prospective weight change in participants of the EPIC-PANACEA study. Am J Clin Nutr 2010;92:398-407. doi:10.3945/ ajcn.2009.28713
- 40 Slimani N, Deharveng G, Unwin I, et al. The EPIC nutrient database project (ENDB): a first attempt to standardize nutrient databases across the 10 European countries participating in the EPIC study. Eur J Clin Nutr 2007;61:1037-56. doi:10.1038/sj.ejcn.1602679
- 41 Slimani N, Kaaks R, Ferrari P, et al. European Prospective Investigation into Cancer and Nutrition (EPIC) calibration study: rationale, design and population characteristics. *Public Health Nutr* 2002;5(6B):1125-45. doi:10.1079/PHN2002395
- 42 Julia C, Touvier M, Méjean C, et al. Development and validation of an individual dietary index based on the British Food Standard Agency nutrient profiling system in a French context. J Nutr 2014;144:2009-17. doi:10.3945/jn.114.199679
- 43 World Health Organization. ICD-10, International Classification of Diseases and Related Health Problems. 10th revision. WHO, 1993.
- 44 Fine JP, Gray RJ. A Proportional Hazards Model for the Subdistribution of a Competing Risk. J Am Stat Assoc 1999;94:496-509. doi:10.108 0/01621459.1999.10474144.
- 45 van Buuren S. Multiple imputation of discrete and continuous data by fully conditional specification. Stat Methods Med Res 2007;16:219-42. doi:10.1177/0962280206074463
- 46 Lange T, Vansteelandt S, Bekaert M. A simple unified approach for estimating natural direct and indirect effects. Am J Epidemiol 2012;176:190-5. doi:10.1093/aje/kwr525
- 47 Gunter MJ, Murphy N, Cross AJ, et al. Coffee Drinking and Mortality in 10 European Countries: A Multinational Cohort Study. Ann Intern Med 2017;167:236-47. doi:10.7326/M16-2945
- 48 Mullee A, Romaguera D, Pearson-Stuttard J, et al. Association Between Soft Drink Consumption and Mortality in 10 European Countries. JAMA Intern Med 2019;e192478. doi:10.1001/ jamainternmed.2019.2478

- 49 Mathur MB, Ding P, Riddell CA, VanderWeele TJ. Web Site and R Package for Computing E-values. *Epidemiology* 2018;29:e45-7. doi:10.1097/FDF.000000000000864
- 50 VanderWeele TJ, Ding P. Sensitivity Analysis in Observational Research: Introducing the E-Value. Ann Intern Med 2017;167:268-74. doi:10.7326/M16-2607
- 51 Masset G, Scarborough P, Rayner M, Mishra G, Brunner EJ. Can nutrient profiling help to identify foods which diet variety should be encouraged? Results from the Whitehall II cohort. Br J Nutr 2015;113:1800-9. doi:10.1017/S000711451500094X
- 52 Mytton OT, Forouhi NG, Scarborough P, et al. Association between intake of less-healthy foods defined by the United Kingdom's nutrient profile model and cardiovascular disease: A population-based cohort study. *PLoS Med* 2018;15:e1002484. doi:10.1371/journal. pmed 1002484
- 53 Chiuve SE, Sampson L, Willett WC. The association between a nutritional quality index and risk of chronic disease. Am J Prev Med 2011;40:505-13. doi:10.1016/j.amepre.2010.11.022
- 54 Streppel MT, Sluik D, van Yperen JF, et al. Nutrient-rich foods, cardiovascular diseases and all-cause mortality: the Rotterdam study. Eur J Clin Nutr 2014;68:741-7. doi:10.1038/ejcn.2014.35
- 55 Willett W, Rockström J, Loken B, et al. Food in the Anthropocene: the EAT-Lancet Commission on healthy diets from sustainable food systems. *Lancet* 2019;393:447-92. doi:10.1016/S0140-6736(18)31788-4
- 56 Scoditti É, Massaro M, Garbarino S, Toraldo DM. Role of Diet in Chronic Obstructive Pulmonary Disease Prevention and Treatment Nutrients 2019:11:1357. doi:10.3390/nu11061357
- 57 Varraso R, Camargo CAJr. Novel dietary risk factors for asthma. Expert Rev Respir Med 2019;13:695-8. doi:10.1080/17476348.2019.16 26721
- 58 Beasley R, Semprini A, Mitchell EA. Risk factors for asthma: is prevention possible?*Lancet* 2015;386:1075-85. doi:10.1016/ S0140-6736(15)00156-7
- 59 Li Z, Rava M, Bédard A, et al. Cured meat intake is associated with worsening asthma symptoms. *Thorax* 2017;72:206-12. doi:10.1136/thoraxjnl-2016-208375
- 60 Varraso R, Chiuve SE, Fung TT, et al. Alternate Healthy Eating Index 2010 and risk of chronic obstructive pulmonary disease among US women and men: prospective study. *BMJ* 2015;350:h286. doi:10.1136/bmj.h286
- 61 Andrianasolo RM, Hercberg S, Kesse-Guyot E, et al. Association between dietary fibre intake and asthma (symptoms and control):

- results from the French national e-cohort NutriNet-Santé. *Br J Nutr* 2019;122:1040-51. doi:10.1017/S0007114519001843
- 62 Zhuang P, Zhang Y, He W, et al. Dietary Fats in Relation to Total and Cause-Specific Mortality in a Prospective Cohort of 521120 Individuals With 16 Years of Follow-Up. Circ Res 2019;124:757-68. doi:10.1161/CIRCRESAHA.118.314038
- 63 Neelakantan N, Koh W-P, Yuan J-M, van Dam RM. Diet-Quality Indexes Are Associated with a Lower Risk of Cardiovascular, Respiratory, and All-Cause Mortality among Chinese Adults. J Nutr 2018;148:1323-32. doi:10.1093/jn/nxy094
- 64 Julia C, Méjean C, Touvier M, et al. Validation of the FSA nutrient profiling system dietary index in French adults-findings from SUVIMAX study. Eur J Nutr 2016;55:1901-10. doi:10.1007/s00394-015-1006-y
- 65 Slimani N, Deharveng G, Southgate DAT, et al. Contribution of highly industrially processed foods to the nutrient intakes and patterns of middle-aged populations in the European Prospective Investigation into Cancer and Nutrition study. Eur J Clin Nutr 2009;63 (Suppl 4):5206-25. doi:10.1038/ejcn.2009.82
- 66 Slimani N, Fahey M, Welch AA, et al. Diversity of dietary patterns observed in the European Prospective Investigation into Cancer and Nutrition (EPIC) project. *Public Health Nutr* 2002;5(6B):1311-28. doi:10.1079/PHN2002407
- 67 Rothman K, Greenland S, Lash T. *Modern Epidemiology.* 3rd ed. Lippincott Williams & Wilkins. 2008.
- 68 Jurek AM, Greenland S, Maldonado G, Church TR. Proper interpretation of non-differential misclassification effects: expectations vs observations. *Int J Epidemiol* 2005;34:680-7 doi:10.1093/ije/dyi060
- 69 Willett WC. Nutritional Epidemiology. 2nd ed. Oxford University Press, 1998. doi:10.1093/acprof:oso/9780195122978.001.0001
- 70 Dubois P, Albuquerque P, Allais O, et al. Effects of front-of-pack labels on the nutritional quality of supermarket food purchases: evidence from a large-scale randomized controlled trial. J Acad Mark Sci 2020; published online 24 April. doi:10.1007/s11747-020-00723-5

**Supplementary information:** FSAm-NPS and Nutri-Score computation, participants flowchart, and additional analyses