Pro-inflammatory dietary intake as a risk factor for CVD in men: a 5-year longitudinal study

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Abstract

Convincing evidence has identified inflammation as an initiator of atherosclerosis, underpinning CVD. We investigated (i) whether dietary inflammation, as measured by the 'dietary inflammatory index (DII)', was predictive of 5-year CVD in men and (ii) its predictive ability compared with that of SFA intake alone. The sample consisted of 1363 men enrolled in the Geelong Osteoporosis Study who completed an FFQ at baseline (2001–2006) (excluding participants who were identified as having previous CVD). DII scores were computed from participants' reported intakes of carbohydrate, micronutrients and glycaemic load. DII scores were dichotomised into a pro-inflammatory diet (positive values) or an anti-inflammatory diet (negative values). The primary outcome was a formal diagnosis of CVD resulting in hospitalisation over the 5-year study period. In total, seventy-six events were observed during the 5-year follow-up period. Men with a pro-inflammatory diet at baseline were twice as likely to experience a CVD event over the study period (OR 2·07; 95% CI 1·20, 3·55). This association held following adjustment for traditional CVD risk factors and total energy intake (adjusted OR 2·00; 95% CI 1·03, 3·96). This effect appeared to be stronger with the inclusion of an age-by-DII score interaction. In contrast, SFA intake alone did not predict 5-year CVD events after adjustment for covariates (adjusted OR 1·40; 95% CI 0·73, 2·70). We conclude that an association exists between a pro-inflammatory diet and CVD in Australian men. CVD clinical guidelines and public health recommendations may have to expand to include dietary patterns in the context of vascular inflammation.

Key words: Dietary inflammatory index: CVD: Risk factors: Longitudinal studies: Nutrition

CVD is the leading cause of death in men in many Western nations including the $USA^{(1)}$ and the $UK^{(2)}$. In Australia, it accounts for one-third of all deaths and is now the leading contributor to direct health care expenditure^(3,4). Although notable improvements in survival rates have been observed

over recent decades, the number of Australian men who are expected to die from repeat heart attacks is projected to increase by >40% by $2020^{(5)}$. These figures highlight the limitations of current primary and secondary prevention strategies. As 80% of the CVD-related burden is attributable to

Abbreviations: DII, dietary inflammatory index; GOS, Geelong Osteoporosis Study.

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modifiable risk factors⁽⁶⁾, there is great potential for health promotion and economic savings through the evaluation of alternative prevention strategies.

Modern-day, obesogenic environments have played a crucial role in propelling the CVD epidemic. Specifically, poor diet quality has been strongly and repeatedly linked to mortality⁽⁷⁾. Indeed, dietary intake has long been recognised as a potent contributor to the pathogenesis of CVD and has received substantial attention⁽⁸⁻¹⁰⁾. Although most of the studies and public health focus regarding CHD have been on its relationship with the intake of individual macronutrients, particularly SFA, new data have highlighted the deleterious effects of inflammationinducing foods that are often used to replace SFA in Western diets (e.g. highly processed carbohydrates, sugar, trans fats, etc.)^(11,12). Accumulating data suggest that CVD is a set of inflammatory disorders, with key processes now thought to lie with the oxidation of LDL-cholesterol and phospholipids. The consequent autoimmune responses are directed against newly formed epitopes, especially oxidised LDL-cholesterol and other oxidised phospholipids⁽¹³⁾.

To date, cardiovascular epidemiology has primarily examined specific nutrients or foods, population means or nutritional guidelines to assess dietary composition within CVD populations, thus neglecting both the potential of dietary-related inflammation and its relation to measures of inflammatory and oxidative stress. It is recognised that a shift away from assessing individual macronutrients or micronutrients towards an investigation of overall diet quality in the pathogenesis of disease is required^(14,15). The utility of novel and innovative methodologies has, thus, been called for in assessing dietary composition as a determinant of disease⁽¹⁶⁾. One such new methodology focuses on the components of diet that are empirically related to markers of immune processes and systemic inflammation in humans. 'Diet-related inflammation' refers to the inflammatory potential of the diet; this is a concept that has been recently utilised in other disciplines such as nutritional psychiatry⁽¹⁷⁾. A pro-inflammatory diet has been linked to a range of somatic conditions and, most recently, indices of obesity⁽¹⁸⁾. We, therefore, aimed to examine the extent to which diet-related inflammation is a risk factor for 5-year cardiovascular incidence in males, using a dietary inflammatory index (DII). We further aimed to compare its predictive ability with that of SFA intake alone.

Methods

Study sample

Details of the Geelong Osteoporosis Study (GOS) have been published elsewhere⁽¹⁹⁾. In brief, GOS comprises an age-stratified, population-based sample of women and men, randomly selected from electoral rolls of the Barwon Statistical Division in south-eastern Australia. As voting is compulsory in Australia for adults aged \geq 18 years, this sampling technique provides a random sample of citizens registered with the Australian Electoral Commission. Population characteristics of this Division are comparable with national levels. Individuals randomly selected from the electoral roll were mailed an invitation letter,

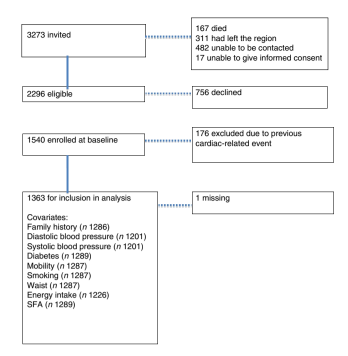


Fig. 1. Recruitment flow chart of Geelong Osteoporosis Study.

with a request to contact the research centre. Those residing in the area for <6 months or those unable to provide informed consent were excluded. In 2001–2006, 3273 men (aged 20–97 years) were invited to participate in the GOS. Of these, 167 had died, 311 had left the region, 482 were unable to be contacted and seventeen were unable to provide informed consent. Of the 2296 remaining individuals, 756 declined to participate (most common reasons: personal 44%, time constraints 26%), leaving 1540 male participants at baseline (participation rate 67%). Fig. 1 displays the recruitment flow chart, representing those included in this study.

Ethics

This study was conducted according to the guidelines laid down in the Declaration of Helsinki, and all the procedures involving human subjects/patients were approved by the Barwon Health Human Research Ethics Committee. Written informed consent was obtained from all the subjects.

Procedure

Although the GOS study comprises ongoing, regular health assessments at the study centre based at Barwon Health (University Hospital Geelong), this study utilised nutrition, anthropometric, demographic and other (non-CVD) health data from GOS baseline assessments. Trained research assistants collected these data during face-to-face interviews with clinical assessments of the participants. Following an overnight fast, blood samples were collected from the participants at the time of baseline assessment at a local pathology laboratory and were stored until batch analysis. In 2011, CVD events data were extracted retrospectively from hospital medical records.

Study measurements

Outcomes. The primary outcome was the occurrence of a CVD event that resulted in hospital presentation over the 5-year follow-up period (post-baseline assessment), with a formal diagnosis of the following: cardiac death, non-fatal myocardial infarction (MI; 'heart attack') based on troponin levels and electrocardiogram readings (ST segment elevation MI or non-ST-segment elevation myocardial infarction (STEMI)), coronary intervention or percutaneous coronary intervention, coronary artery bypass grafting, amputation or transient ischaemic attack, ischaemic or haemorrhagic stroke. Data were extracted from catchment area hospital medical records held at Barwon Health by medically trained research fellows. This procedure was likely to capture the majority of CVD events in the region due to three key reasons: (i) Barwon Health was the sole national health service emergency facility in the regional catchment area at the time, (ii) admission was accessible to the general public with no out-of-pocket expense and (iii) only a small percentage of participants moved away from the Barwon region during follow-up. Participants gave consent for their hospital admission records to be accessed, allowing CVD events data to be recorded for all the participants, regardless of retention status at subsequent followup assessments.

Exposure: dietary inflammatory index. Details of the development of the DII have been documented elsewhere⁽²⁰⁾, including its validation against general inflammatory biomarkers⁽²¹⁾. Data were derived from Victorian Cancer Council FFQ completed by the participants at baseline used to assess participants' usual consumption of seventy-four foods and six alcoholic beverages over the previous year on a ten-point frequency scale. Raw data provided detailed information about the consumption of common foods (fruit, vegetables, dairy products, meat, fish, pasta, rice, breads, nuts, legumes, breakfast foods, snack foods and alcohol). The FFQ provided estimated intakes of total carbohydrates and specific macronutrients (e.g. SFA, MUFA, PUFA, proteins, carbohydrates, sugars, starch, dextrins, fibre, alcohol and fatty acids), a range of micronutrients and glycaemic load.

DII scores were computed from these FFQ data by a trained research fellow. Using a regionally representative world database (food consumption from eleven populations globally), means and standard deviations were calculated for each parameter. The 'standard mean' was subtracted from the actual exposure and divided by its standard deviation. These Z scores were converted to percentiles (minimising effects of outliers/right-skewing)⁽²⁰⁾ and centred by doubling the value and subtracting 1 to achieve symmetrical distribution of the scores. The product for each food parameter and the adjusted article score was calculated and summed across the food parameters to generate an overall DII score for each participant⁽²⁰⁾, where higher values indicate a more pro-inflammatory diet. The steps to calculate DII are shown in Fig. 2. Overall, twenty-two of the forty-five possible food parameters used for DII calculation were available in this study, and those food parameters were carbohydrates, proteins, total fat, energy, fibre, cholesterol, SFA, MUFA, PUFA, n-3, n-6, niacin,

thiamine riboflavin, Fe, Mn, Zn, vitamin A, vitamin E, folic acid and β -carotene.

Covariates. Data on family history of CVD, age, waist circumference, smoking status, frequency of alcohol use, mobility levels, systolic and diastolic blood pressures (BP), energy intake and diabetes status were collected at baseline assessment. Body weight (sp 0.1 kg) was measured using electronic scales, and height (sp 0.1 cm) was measured using a wall-mounted stadiometer (Harpenden). Waist circumference was measured with a tape measure around the smallest circumference between the lower rib and iliac crest, and BP (seated) was measured using a digital meter. Data on smoking status, alcohol use and exposure to drugs were obtained through self-report. Daily energy intake (kJ/d) was derived from the FFQ. Blood samples were stored in serum aliquots at -80°C from which blood glucose levels were calculated using standard laboratory procedures. For the purpose of this study, diabetes diagnosis at baseline was based on at least one of the following: a blood glucose reading of $\geq 7.0 \text{ mmol/l}$, use of antihyperglycaemic medication or self-report at baseline assessment. SFA consumption (mg/d) was computed from the dietary data by means of the nutrient composition tables in the NUT-TAB95 database (Food Standards Australia New Zealand, Canberra, 1995).

Data analysis

Descriptive statistics were used to identify the key characteristics of the sample. As our *a priori* hypothesis was that a pro-inflammatory diet would be associated with greater odds of a CHD event when compared with (what has been shown to be a cardio-protective diet) an anti-inflammatory dietary intake, DII scores were dichotomised such that positive values represented more pro-inflammatory diets and negative scores reflected more anti-inflammatory diets. Using methods described by Hosmer & Lemeshow⁽²²⁾, logistic regression was performed to examine the predictive effect of DII grouping at baseline (where anti-inflammatory dietary intake was treated as the referent group) on the primary outcome of the occurrence of a 5-year CVD event.

Logistic regression was used, as the outcome of interest was not common in our study population (<10%); seventy-six events were recorded out of 1363 participants. Further, as all the participants were followed-up to the outcome of interest (CVD admission data were available from 1993), we explored the time to event, ignoring censoring.

Participants with an admission for a CVD event (as defined by the primary outcome) at any time before study enrolment were identified and subsequently excluded (n 176) from the analyses in order to determine the directionality of the relationship of interest, leaving a sample size of 1363 for inclusion in the statistical models. Missing data for each covariate are displayed in Fig. 1. We were particularly interested in the magnitude of effect of SFA on CVD incidence, relative to the DII, given that it has been considered a putative risk factor for atherosclerotic progression. As this macronutrient was a constituent of the DII, we estimated its effect in an independent model, in order to

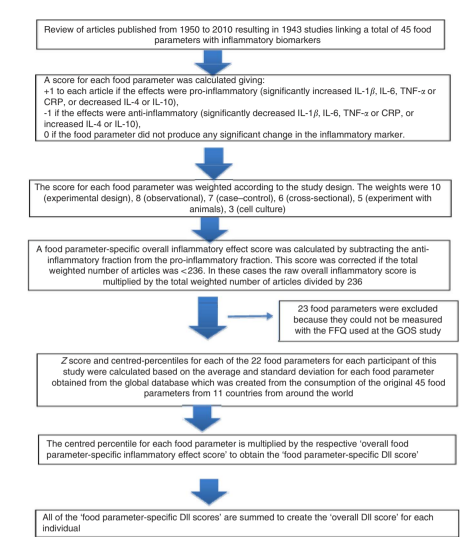


Fig. 2. Sequence of steps in creating the dietary inflammatory index (DII) in the Geelong Osteoporosis Study (GOS). CRP, C-reactive protein.

prevent problems with inter-collinearity and to enhance model stability. Logistic regression was also used to examine the relationship of SFA intake with CVD incidence by applying the same technique, but substituting the independent variable, DII score, for recommended daily SFA intake⁽²³⁾ (<25 g/d).

We first conducted a univariate analysis to examine the unadjusted association between the independent and dependent variables. The second iteration of the model adjusted for the traditional CVD risk factors of age, diabetes, systolic and diastolic BP, smoking history, activity level (sedentary v. active) and waist circumference, as well as total daily energy consumption. Alcohol intake was excluded from the statistical models as it forms a component of the DII. A sensitivity analysis was performed excluding those men who indicated that they were using hypocholesteraemic agents at baseline. OR and accompanying 95 % CI were presented for each model. Cohen's d effect sizes were reported for each model, calculated from the R^2 values generated. As a secondary outcome, a survival analysis was conducted to explore the effect of pro- and anti-inflammatory dietary patterns on time to first CVD event

over the study period (years), presented as a Kaplan–Meier plot. All the analyses were performed using $Stata^{(\!\!\!\!\ensuremath{\mathbb{R}})}$ version 13.

Results

Characteristics of sample

Key characteristics of the sample are shown in Table 1. Compared with those recording a more anti-inflammatory diet, participants with a more pro-inflammatory dietary intake were older and had high BP. A larger proportion of participants in this group was sedentary, smokers, used aspirin or statins, had diabetes and had a family history of CVD. With respect to nutritional profile, this group had lower daily macronutrient intake (overall, SFA, PUFA and MUFA, protein and carbohydrates), lower intake of all micronutrients and a higher glycaemic index (Table 1). In total, 131 men were using anti-inflammatory drugs at baseline. The overall range of DII scores for the sample was –2:96 to 2:63. Mean baseline DII scores for the two groups were 1:00 and

A. O'Neil et al.

Table 1. Key characteristics of the sample by baseline dietary inflammation status (Mean values and standard deviations; percentages and 95 % confidence intervals; *n* 1363)

Variables	Anti-inflammatory diet		Pro-inflammatory diet	
	Mean	SD	Mean	SD
Age (years)	53-2	19.2	59.0	19.2
Height (cm)	177.5	34.6	175.6	41.9
Waist (cm)	101.1	65.6	103.7	72.8
Systolic blood pressure (mmHg)	141.4	90.2	149.9	121-4
Diastolic blood pressure (mmHg)	87.7	94.8	97.3	128.0
Regular alcohol use (weekly intake)	20.0	21.9	19.5	22.2
Nutritional energy intake (kJ)	10 896.9	3378.5	7130.4	1874.9
Fat intake (g)	109.5	43.0	71.1	23.4
SFA intake (g)	44.0	19.8	29.1	11.0
PUFA (g)	17.1	7.0	11.2	5.2
MUFA (g)	39.0	16.2	24.8	8.6
Protein intake (g)	125.4	46.9	78.9	23.3
Carbohydrate intake (g)	281.9	81.7	189.4	53.6
Sugar intake (g)	122.6	40.4	82.8	29.8
Starch intake (g)	157.9	54.5	105.7	36.5
Fibre intake (g)	31.0	9.8	18.5	5.6
β -Carotene intake (µg)	3923.6	1595.0	2090.7	1007.3
Ca intake (mg)	1124.9	358-0	814·0	297.4
Dietary cholesterol intake (mg)	390.6	169-0	251.1	99.2
Folate intake (µg)	388.4	117.6	230.6	68·5
Fe intake (mg)	18.8	6.8	11.0	3.30
Mn intake (mg)	395.9	118-6	244.5	64·7
Niacin intake (mg)	31.5	12.0	17.9	5.9
P intake (mg)	2040-4	629.5	1313.4	363.7
K intake (mg)	3870.8	979.1	2436.4	616-5
Retinol intake (µg)	518-3	213.9	378.8	162.7
Riboflavin intake (mg)	3.3	1.3	2.0	0.7
Na intake (mg)	3613.9	1320.6	2327.8	737.6
Thiamin intake (mg)	2.3	0.8	1.4	0.5
Vitamin C intake (mg)	179.5	83.1	97·0	51·8
Vitamin E intake (mg)	9.14	2.7	5.6	1.8
Zn intake (mg)	16.5	6.7	10.1	3.2
	53.5	3.9	54.9	3·2 4·7
Glycaemic index Baseline DII score	-1.0	0.7	1.0	4·7 0·7
5-year DII score Time to first CVD event (years)	-0·4 2·7	1·1 1·3	0.7 2.6	1.0 1.3
Time to mat OVD event (years)				-
	%	95 % CI	%	95 % CI
Family history of CVD (yes)	49.7	45.7, 53.8	50.3	46.9, 53
Depression (current, major)	11.1	11.7, 56.8	9.1	0.9, 50.0
Statin use (yes)	11.6	9.2, 14.4	19.0	16.5, 21.9
Aspirin use (yes)	10.7	8.5, 13.5	16.7	14.3, 19.4
Smoking status (ever smoked)	48.9	44.9, 52.9	57.3	0.54, 60
Mobility level (sedentary)	20.1	17.1, 23.5	27.3	24.3, 30.
Diabetes (yes)	6.4	4.7, 8.6	7.7	6.1, 9.8

DII, dietary inflammatory index.

-1.00, respectively. In total, seventy-six events were observed during the 5-year follow-up period.

Impact of baseline dietary inflammatory index scores on the presence of 5-year CVD events

Univariate analyses revealed that men recording a pro-inflammatory diet were twice as likely to experience a CVD event over the 5-year observation period when compared with those consuming an anti-inflammatory diet (Table 2). Importantly, this association held after adjustment for traditional CVD risk factors and total energy intake (effect size d = 0.20). As older age exerted a strong effect within the main effects model, we subsequently explored whether age was an effect modifier using an age-by-baseline DII

score interaction variable. Although its inclusion strengthened the relationship between DII and CVD occurrence, the interaction effect was not 'statistically significant' (data not shown).

Finally, we adjusted for usage of medications with antiinflammatory properties (e.g. aspirins, statins). The inclusion of this variable into the final model did somewhat attenuate the association between a pro-inflammatory diet and CHD (OR 1.95; 95 % CI 0.98, 3.87).

Relationship between baseline daily SFA intake and presence of 5-year CVD events

As a *post hoc* analysis, we were interested in the magnitude of the relationship between a pro-inflammatory diet and CVD Table 2. Baseline dietary inflammatory index (DII) category (pro-inflammatory v. anti-inflammatory) as a predictor of 5-year cardiovascular event (presence/absence)

(Odds ratios and 95% confidence intervals)

Variables	Unadjusted		Adjusted (model 1)	
	OR	95 % CI	OR	95 % CI
Pro-inflammatory diet (as estimated by DII score)	2.07*	1.20, 3.55	2.00*	1.01, 3.96
Family history of CVD (yes)	_		2.75*	1.55, 4.93
Systolic blood pressure (mmHg)	_		1.00	0.99, 1.02
Diastolic blood pressure (mmHg)	_		0.99	0.98, 1.01
Mobility (sedentary)	_		0.97	0.62, 1.49
Diabetes status (no)	_		0.70	0.29, 1.70
Smoking history (never)	_		0.86	0.49, 1.50
Waist circumference	_		1.00	0.97, 1.02
Age (years)	_		1.08*	1.06, 1.10
Total daily energy intake (kJ)	_		1.00	0.99, 1.00

* P<0.05 level.

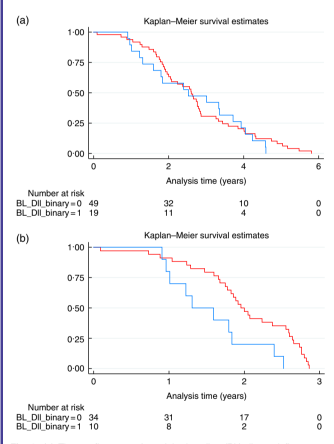


Fig. 3. (a) Time to first event (years) by baseline (BL) dietary inflammatory index (DII) grouping. (b) Time to first event (0–3 years) by baseline DII grouping.

——, Pro-inflammatory diet; ——, anti-inflammatory diet.

events as it compared with that of SFA intake. Univariate analyses revealed that high SFA intake significantly predicted the occurrence of 5-year CVD events (OR 1.87; 95% CI 1.12, 3.10); however, this relationship was no longer significant after the inclusion of traditional CVD risk factors (diabetes and lipid markers were excluded to avoid over-fitting the model) (adjusted OR 1.74; 95% CI 0.92, 3.26). There were no significant interactions between DII scores and SFA scores.

Impact of dietary inflammatory index scores on time to first CVD event

The association between DII group and time to first CVD event was also investigated. Using a likelihood ratio test, the model met the proportional hazards assumption. The seventy-six observations comprised 204.5 years total analysis time at risk and under observation. Kaplan–Meier curves are displayed for the overall sample for the study period (Fig. 3a) and for the events occurring during the first 3 years after enrolment (Fig. 3b, in which it appeared that the inter-group differences were most pronounced).

Discussion

In a longitudinal, nationally representative sample of Australian men, a pro-inflammatory diet (as estimated by the DII) was a significant predictor of CVD occurrence over 5 years. When compared with those consuming an anti-inflammatory diet, this dietary pattern more than doubled the likelihood of a CVD event that resulted in hospitalisation, even after adjustment for traditional CVD risk factors. The use of anti-inflammatory medication, however, did attenuate the observed relationships. The time to first CVD event did not significantly differ by dietary intake over the 5-year study period. When we examined the specific relationship of daily SFA consumption to 5-year CVD incidence, a non-statistically significant association was observed.

To date, evidence that a diet primarily characterised by its pro-inflammatory properties is a potential risk factor for CVD has been lacking. Although other studies have suggested that dietary patterns of inflammatory nature may be linked to heart disease⁽²⁴⁾, to our knowledge, this is the first study to investigate and, moreover, demonstrate the ability of the DII to predict CVD events. Our findings are intuitively consistent with results of epidemiological studies highlighting the benefits of an anti-inflammatory diet. The Mediterranean diet score, for example, has been shown to predict CVD mortality in Australian men with diabetes in whom a 6% reduction in risk per unit of Mediterranean diet score (adjusted hazard ratio (HR) 0.94; 95% CI 0.89, 0.99) was observed⁽²¹⁾. Similar effects have been

observed under real-world, randomised conditions. PREDIMED investigators demonstrated that a Mediterranean diet supplemented with extra-virgin olive oil (adjusted HR 0·70; 95% CI 0·54, 0·92) or nuts (adjusted HR 0·72; 95% CI 0·54, 0·96), when compared with a low-fat diet, led to a reduction in the incidence of major cardiovascular events in those with high CVD risk⁽²⁵⁾. In that study, the DII was associated with higher average BMI, waist circumference and waist:height ratio after adjusting for known risk factors and remained associated with obesity after controlling for the effect that adherence to Mediterranean diet score had on inflammation⁽¹⁸⁾.

Our secondary finding suggests that the combined effect of inflammatory dietary constituents on the occurrence of CVD could be more potent than the effects of SFA alone. Traditionally, the impact of pro-inflammatory dietary patterns on disease onset has been somewhat obscured by the attention paid to putative, individual dietary risk factors such as SFA. Esposito & Giugliano⁽²⁶⁾ and others have provided evidence of the pro-inflammatory milieu as a mechanism through which unhealthy diets may induce cardio-metabolic diseases. They recommend that fully explicating the link between diet and inflammation could 'elucidate the mechanisms by which dietary patterns improve cardiovascular health'. This study could not elucidate the mechanisms underpinning the relationship between diet, inflammation and CVD and, importantly, did not have the required lipid profile data. However, it provides preliminary evidence to support a movement towards viewing dietary intake in terms of totality of effect and not just as a source of specific culprit nutrients. Although SFA intake is part of the DII, which was associated with CVD outcomes, in isolation SFA was not strongly associated with these outcomes and, by inference, the atherosclerotic state.

Data from animal models, population-based studies and randomised trials provide some support for the key role of dietary intake on systemic inflammation. Epidemiological evidence has linked general markers of inflammation – including C-reactive protein, IL-6 and TNF – to greater consumption of 'Western-style' diets, which are high in sweetened soft drinks, refined grains, red meat, margarine and fish, but low in wine, coffee, olive oil, and green leafy and yellow vegetables⁽²⁷⁾. A sucrose-rich diet in overweight adults over 10 weeks resulted in significant increases in inflammatory markers (haptoglobin and transferrin) and small increases in C-reactive protein levels⁽²⁸⁾. There is preliminary evidence of dysfunction of immune system activation induced by diet via the gut microbiota⁽²⁹⁾.

Overall, the diet of the group consuming a more proinflammatory diet contained a range of nutritional deficiencies when compared with those consuming a more anti-inflammatory diet. Despite the latter group appearing to consume higher levels of macronutrients that could be considered inflammatory (e.g. *SFA intake, sugar intake*), closer examination of the characteristics of the anti-inflammatory group's diet (at a micronutrient level) suggests that it is much more nutrient dense with phytochemicals and micronutrients commonly found in vegetables and fruits. For example, elevated levels of β -carotene, folate and vitamin C suggest higher fruit and vegetable intake, whereas low glycaemic intake indicates that whole grains are key constituents of their carbohydrate consumption. It is plausible that the nutritionally superior profile of the anti-inflammatory group is contributing to better overall health and well-being.

Strengths and limitations

The key strength of this study was that it is the first to explore the utility of a novel, DII as a key marker of CVD. To date, the DII has been found to be associated with inflammatory cytokines including C-reactive protein and IL-6(30), increased odds of asthma and reduced lung function in an Australian population⁽³¹⁾, shiftwork⁽³²⁾, colorectal cancer among women in the Iowa Women's Health Study⁽³³⁾, prostate cancer⁽³⁴⁾ and pancreatic cancer⁽³⁵⁾ in case-control studies in Italy. The longitudinal study design, nationally representative sample, use of comprehensive nutritional data (to both generate DII scores and inform statistical modelling), as well as the use of 'hard' clinical end points are additional strengths. Our primary finding adds to the burgeoning evidence base that dietary inflammation is associated with other chronic conditions. The major limitation of the study was the absence of lipid profile data for inclusion in our statistical models (for which other key risk factors for CVD were adjusted). Second, the absence of robust physical activity data is a limitation of the study. However, the inclusion of a proxy marker (mobility) in the adjusted model was not a significant predictor of CVD outcome.

A final limitation of the study was the likely lack of power to detect time to event differences over the study period due to the small number of cases. Future studies with adequate power to detect such differences between dietary inflammation groups and time to event should be carried out.

Conclusions

In conclusion, this study could be an important first step in broadening our understanding of the contribution of dietary intake to CVD risk to include dietary-promoted inflammation. Although there are numerous models of dietary interventions targeting CVD prevention (nutrient-based; e.g. Heart Foundation, Mediterranean diet), there is likely to be potential benefits associated with applying an approach to CVD prevention that is based on intricate inflammatory biological pathways from the whole diet. This moves away from a reductionist approach that depends on identifying specific nutrients or foods as risk factors, but rather identifies a cardio-protective diet that is independent of any pre-determined nutrient or cuisine models, population means or nutritional guidelines. Studies are now required to confirm these findings as well as those that can quantify the extent to which reducing dietary inflammation has health and economic benefits in the long term. Following recent calls, other future studies looking at this type of model on long-term CVD-related mortality are required⁽³⁶⁾.

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A. O'N. and F. N. J. designed the research; J. A. P. and M. A. K. oversaw all the aspects of the GOS study; K. K. collected the CVD data; F. N. J. coordinated the collection of dietary data; J. R. H. developed the DII instrument and DII scores were generated under his supervision by N. S.; and A. O'N. analysed the data, wrote the manuscript and had primary responsibility for its final content. All the authors read and approved the final version of the manuscript.

None of the authors has any conflicts of interest with the NHMRC.

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