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# The association between dietary patterns and the novel inflammatory markers platelet-activating factor and lipoprotein-associated phospholipase A<sub>2</sub>: a systematic review

Carolyn J. English, Hannah L. Mayr, Anna E. Lohning, and Dianne P. Reidlinger 💿

#### 12

**Context:** Atherosclerosis is a disease of chronic inflammation. Recent research has identified 2 novel inflammatory biomarkers: platelet-activating factor (PAF) and lipoprotein-associated phospholipase  $A_2$  (Lp-PLA<sub>2</sub>). Diet has been proposed as a mediator of inflammation, but to date, the focus for these novel biomarkers has been on individual foods and nutrients rather than overall dietary patterns. **Objective:** To systematically review the literature on the association between dietary patterns and PAF and Lp-PLA<sub>2</sub>. Data Sources: The PubMed, Embase, CINAHL, and Cochrane CENTRAL literature databases were searched. Data Analysis: Study quality was evaluated using the Quality Criteria Checklist. Sixteen studies (n = 4 observational and n = 12 interventional) were included and assessed for associations between dietary patterns and PAF and Lp-PLA<sub>2</sub>. Conclusion: Study quality varied from neutral (n = 10) to positive (n = 6). Mediterranean, heart healthy, and vegetarian dietary patterns were associated with improved levels of PAF and Lp-PLA<sub>2</sub>. Conversely, Western dietary patterns were less favorable. A range of wellestablished, healthier dietary patterns may lower inflammation and the risk of atherosclerosis. More well-designed studies are needed to confirm these findings and identify other dietary patterns that improve inflammation.

# INTRODUCTION

Atherosclerosis, the main underlying cause of cardiovascular disease (CVD), is a chronic arterial disease leading to fatty streaks and atheromas in the arterial wall.<sup>1,2</sup> Once thought to be solely caused by dyslipidemia, atherosclerosis is now known to be a result of inflammatory responses.<sup>3</sup> Inflammation is involved in all stages of atherosclerosis, from the initial injury of the endothelium to plaque formation and eventual plaque rupture and thrombosis.<sup>4,5</sup>

Two novel inflammatory markers involved in CVD that are receiving increasing attention are plateletactivating factor (PAF) and lipoprotein-associated phospholipase  $A_2$  (Lp-PLA<sub>2</sub>).<sup>6,7</sup> PAF is the most potent lipid inflammatory mediator and is produced upon

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Key words: cardiovascular disease, dietary patterns, inflammation, lipoprotein-associated phospholipase A<sub>2</sub>, Lp-PLA<sub>2</sub>, PAF, platelet-activating factor.

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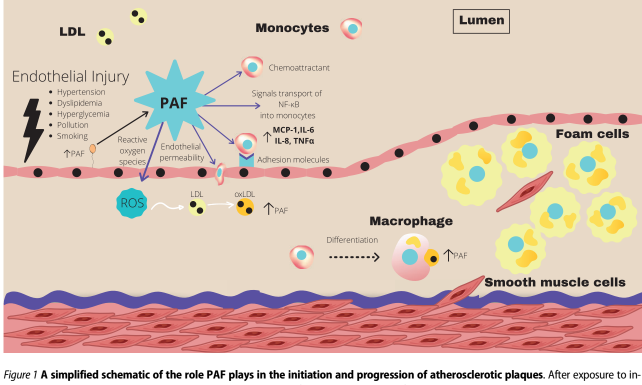
This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs licence (https:// creativecommons.org/licenses/by-nc-nd/4.0/), which permits non-commercial reproduction and distribution of the work, in any medium, provided the original work is not altered or transformed in any way, and that the work is properly cited. For commercial re-use, please contact journals.permissions@oup.com stimulation by numerous cells such as platelets, endothelial cells, and leukocytes.<sup>8,9</sup> PAF is implicated in every step of atherosclerosis (Figure 1).<sup>4,6,10,11</sup> PAF plays a crucial role in the initiation of atherosclerosis and one of its main pro-inflammatory actions is the mediation of adhesion of monocytes to the endothelium and initiation of gene transcription within monocytes to produce inflammatory cytokines such as monocyte chemoattractant protein-1, interleukin (IL) 8, and tumor necrosis factor  $\alpha$  (TNF- $\alpha$ ).<sup>12,13</sup> PAF also stimulates the release of the proinflammatory cytokine IL-6 from both endothelial cells and monocytes.<sup>14</sup>

PAF induces an influx of Ca<sup>2+</sup>, which results in increased endothelial permeability as the endothelial cells contract, allowing the migration of low-density lipoprotein (LDL) cholesterol and monocytes into the intima.<sup>15–18</sup> PAF also stimulates reactive oxygen and nitrogen species and contributes to the oxidation of LDL.<sup>6,19</sup> PAF is further involved in the differentiation of monocytes into pro-inflammatory macrophages that engulf oxidized LDL, and is involved in the formation of foam cells and the growth and rupture of plaques.<sup>20,21</sup>

PAF, once produced, triggers an uncontrolled and prolonged inflammatory milieu, because it is responsible for the production of new PAF molecules and additional free radicals.<sup>21,22</sup> Patients with diabetes, heart failure, acute myocardial infarction, and coronary heart disease have elevated levels of PAF.<sup>23–28</sup>

Lp-PLA<sub>2</sub> (alternatively known as platelet-activating factor-acetylhydrolase) is an enzyme that catalyzes hydrolysis of PAF and belongs to the PLA<sub>2</sub> superfamily.<sup>29</sup> As Lp-PLA<sub>2</sub> hydrolyses PAF into the inactive form lyso-PAF, Lp-PLA<sub>2</sub> levels are proposed to be determined by in vivo levels of PAF and may serve as a reliable surrogate marker of PAF.<sup>30</sup> Because Lp-PLA<sub>2</sub> catabolizes PAF, Lp-PLA<sub>2</sub> appears to play an anti-inflammatory role. However, because of its nonspecificity for its ligand, the hydrolysis products of Lp-PLA<sub>2</sub> have been linked to pathologies.<sup>31</sup>

Lp-PLA<sub>2</sub> is primarily secreted by macrophages and circulates in the blood bound to LDL and high-density lipoprotein (HDL), with the majority attached to LDL, and preferentially to small dense fractions.<sup>32</sup> It is proposed that HDL bound to Lp-PLA<sub>2</sub> plays a protective



*Figure 1* **A simplified schematic of the role PAF plays in the initiation and progression of atherosclerotic plaques**. After exposure to injury, the endothelial cell is activated, triggering the production of PAF and expression of adhesion molecules. PAF acts as a strong chemoat-tractant and mediates the firm adhesion of monocytes to the endothelium via adhesion molecules. PAF signals the transport of NF- $\kappa$ B into the nucleus of the monocytes, triggering gene transcription of pro-inflammatory cytokines such as MCP-1, IL-6, IL-8, and TNF- $\alpha$ . PAF stimulates the production of ROS, which contributes to the oxidation of LDL. PAF reduces endothelial nitric oxide production and increases endothelial permeability, allowing the transmigration of LDL and monocytes into the intima. PAF is responsible for the differentiation of monocytes into macrophages that engulf oxLDL, which triggers the production of more PAF. *Abbreviations*: IL, interleukin; NF- $\kappa$ B, nuclear factor  $\kappa$ B; LDL, low-density lipoprotein; MCP-1, monocyte chemoattractant protein-1; oxLDL, oxidized low-density lipoprotein; PAF, platelet-activating factor; ROS, reactive oxygen species; TNF- $\alpha$ , tumor necrosis factor  $\alpha$ .

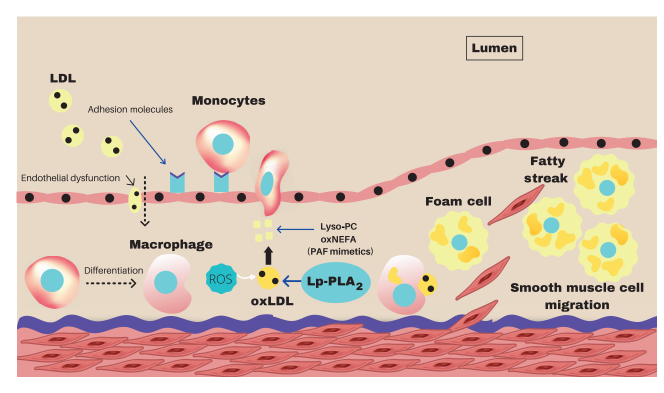
role, whereas LDL-bound Lp-PLA<sub>2</sub> is atherogenic.<sup>32</sup> When associated with LDL, Lp-PLA<sub>2</sub> hydrolyzes oxidized phospholipids on the surface of the LDL particles, creating pro-inflammatory and pro-atherogenic by-products such as lysophosphatidylcholine and oxidized, nonesterified fatty acids.<sup>33</sup> Lysophosphatidylcholine and oxidized, nonesterified fatty acids mimic PAF in mediating inflammation by upregulating adhesion molecules; acting as a chemoattractant to monocytes; activating leukocytes; stimulating cytokine production such as IL-6 and TNF- $\alpha$ ; contributing to necrosis and apoptosis of macrophages in the plaque; and inducing smooth muscle migration into the intima (Figure 2).<sup>31,34–37</sup> Lp-PLA<sub>2</sub> is an independent risk marker for coronary heart disease events, stroke, calcific aortic-valve stenosis, and plaque stability.<sup>38–41</sup>

Previous research on diet and PAF and/or Lp-PLA<sub>2</sub> is limited. However, some research has demonstrated that bioactive compounds found in foods regularly consumed in the traditional Mediterranean diet contain natural PAF inhibitors.<sup>20</sup> These compounds inhibit inflammation by preventing PAF from binding to its receptor, blocking the cascade of intracellular signaling and inflammatory processes, and possibly by inhibiting

metabolic enzymes used in the remodeling pathway for PAF synthesis.<sup>42–44</sup> This research provides some insight into the potential mechanisms of components within the Mediterranean diet and its established cardioprotective effects.<sup>45</sup>

Research into specific Mediterranean foods that inhibit PAF have predominantly been in vitro studies using washed rabbit platelets and, more recently, human platelets.<sup>46</sup> The foods include fish<sup>47,48</sup>; eggs<sup>49</sup>; honey<sup>50</sup>; wild plants<sup>51</sup>; garden peas<sup>52</sup>; dairy (especially fermented and of goat and sheep origin)<sup>53–56</sup>; goat and sheep meat<sup>57</sup>; flaxseeds<sup>58</sup>; olive oil and olive pomace<sup>59–61</sup>; wine<sup>46</sup>; grapes<sup>62</sup>; *Origanum onites* (Cretan oregano)<sup>63</sup>; clove and cinnamon<sup>64</sup>; onion<sup>65</sup>; garlic<sup>66</sup>; and seeds oils, such as corn, sunflower, and sesame.<sup>59</sup> Foods found outside the Mediterranean region that inhibit PAF include soy sauce,<sup>67</sup> *Camillea sinensis* (tea),<sup>68</sup> and curcumin.<sup>69</sup>

Dietary effects on Lp-PLA<sub>2</sub> levels are largely unexplored, but some evidence from studies in humans has shown that low-energy diets with concurrent weight loss can reduce Lp-PLA<sub>2</sub> levels, whereas increased energy intake is associated with higher Lp-PLA<sub>2</sub> levels.<sup>70,71</sup>



*Figure 2* Lp-PLA<sub>2</sub> involvement in the progression of atherosclerosis. Lp-PLA<sub>2</sub> circulates primarily bound to LDL cholesterol, concentrating in small dense LDL. After oxidation of LDL, Lp-PLA<sub>2</sub> hydrolyzes oxLDL, creating 2 inflammatory phospholipids, lyso-PC and oxNEFA, both of which mimic PAF. Lyso-PC and oxNEFA upregulate inflammatory mediators such as adhesion molecules, MCP-1, IL-6, and TNF-*α*; contribute to endothelial dysfunction; promote chemotaxis, drawing monocytes into the arterial intima; trigger smooth muscle cell migration; and induce apoptosis and cytotoxic effects contributing to necrotic core growth. *Abbreviations*: LDL, low-density lipoprotein; Lp-PLA<sub>2</sub>, lipoprotein-associated phospholipase A<sub>2</sub>; Lyso-PC, lysophosphatidylcholine; MCP-1, monocyte chemoattractant protein-1; oxLDL, oxidized low-density lipoprotein; convEFA, oxidized nonesterified fatty acids; PAF, platelet-activating factor; ROS, reactive oxygen species; TNF-*α*, tumor necrosis factor alpha.

The replacement of 5% of energy from carbohydrates with energy from protein is associated with a decrease in Lp-PLA<sub>2</sub> activity.<sup>72</sup> An 8-week intervention with the supplementation of omega-3 fatty acids did not influence Lp-PLA<sub>2</sub> activity in older adults,<sup>73</sup> whereas a similar 30-day intervention in people with stable coronary artery disease resulted in decreased Lp-PLA<sub>2</sub> levels.<sup>74</sup>

Studies have varied in terms of the assays used to measure Lp-PLA<sub>2</sub>. Lp-PLA<sub>2</sub> assays can measure either plasma concentrations or enzymatic activity. This makes comparisons between studies and interpretation of results difficult. Enzyme activity assays now predominate the recent literature, because mass assays have been shown to be less accurate for risk stratification, because of their ability to only detect a smaller amount of Lp-PLA<sub>2</sub>, particularly that associated with HDL.<sup>75,76</sup>

In a recent review considering 17 studies of varying designs that investigated the Mediterranean diet and its components, the authors concluded that this dietary pattern has the potential to lower PAF and Lp-PLA<sub>2</sub> levels.<sup>30</sup> However, the scope of that review was limited to 1 database, and 12 of the 17 included studies examined individual foods, alcohol, or supplements such as fish oil and eicosapentaenoic acid, and not dietary patterns, which are more translatable and relevant across populations. In the present review, we aimed to comprehensively investigate the association between overall dietary patterns and their effect on PAF and Lp-PLA<sub>2</sub> as novel inflammatory biomarkers.

#### MATERIALS AND METHODS

For this systematic review, we followed the requirements of the Preferred Reporting of Systematic Reviews and Meta-Analyses (PRISMA) statement (Supporting Information online), and the review was registered in July 2021 with the International Prospective Register of Systematic Reviews (PROSPERO no. CRD42020169666; available at http://www.crd.york.ac.uk/PROSPERO).

# Search strategy

The databases PubMed, Embase, CINAHL, and Cochrane CENTRAL were searched for relevant studies, with backward citation checking of relevant reviews retrieved in the search. A search for trial protocols through the ClinicalTrials.gov website (www.clinicaltrials.gov) and World Health Organization International Clinical Trials Registry Platform (https://apps.who.int/trialsearch/) was also performed. Databases were searched from inception; the search date was February 21, 2020, with an update to the search performed on February 7, 2021. Table 1 lists PICOS criteria (ie, participants, intervention, comparators, outcomes, and study designs) used to identify studies for inclusion. Eligible studies in any language were considered, provided they were full articles published in a peer-reviewed journal.

A comprehensive search strategy was developed by the research team in conjunction with an experienced librarian. Terms used in the literature search included PAF, platelet-activating factor, Lp-PLA<sub>2</sub>, lipoprotein-associated phospholipase A2, diet, and variations of these terms. The complete search strategy is available in the Supporting Information online.

#### Data management and extraction

Search results were imported into Endnote, version X9.3.3,<sup>77</sup> for de-duplication, then uploaded to Covidence<sup>78</sup> for removal of duplicates and screening. Screening of titles and abstracts against the inclusion criteria was undertaken independently and in duplicate by 2 researchers. Full-text articles were then reviewed independently and in duplicate by 2 researchers and screened for inclusion criteria. Disagreements were resolved by discussion or by a third reviewer.

Data extraction was performed by populating dataextraction tables for multiple study designs from the *Cochrane Handbook for Systematic Reviews of Interventions*,<sup>79</sup> which were further adapted to extract

Table 1 <b>PICOS criteria</b>	for inc	lusion and	exclusion	of stud	ies
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Parameter	Inclusion criteria	Exclusion criteria
Participants	Adults $\geq$ 18 y	Aged $<$ 18 y
Intervention	Studies examining diet assessed by dietary patterns, dietary scores, dietary indices, and food patterns	Studies reporting animal or cellular models, or that analyzed consumption of single nutrients or foods rather than a dietary pattern
Comparator	Any/none	Any/none
Outcome	Any measurement of systemic inflammation using PAF and/or Lp-PLA <sub>2</sub> . Secondary outcomes included other reported novel markers of inflammation	Other cardiovascular disease outcomes
Study design	Observational (eg, prospective cohort, retrospective cohort, cross sectional, longitudinal, case-control, case series), intervention and randomized controlled trials	None

additional information during this stage. Data extraction was piloted on included articles reporting 3 different study designs, and then was amended to a final format. Data extraction was undertaken by 1 researcher and independently reviewed for accuracy by another researcher.

Data extracted included author, date published, study design, level of evidence, population, sex, country, age, type of dietary pattern, control group, sample size, and study duration. Primary outcomes extracted were PAF levels, PAF-induced platelet aggregation in platelet-rich plasma, specific activities of plasma lyso-PAF and PAF-AH, and LP-PLA<sub>2</sub> mass and activity. Secondary outcomes extracted were any reported biomarkers identified as novel (ie, not recognized as a common inflammatory marker by the research team) and related to CVD. Study authors were contacted by email for additional information if required data had not been published.

#### Outcomes

The primary outcomes included mean net change in outcome measurements (ie, blood PAF, lyso-PAF, and PAF-AH levels; Lp-PLA<sub>2</sub> mass and/or activity; or plate-let aggregation induced by PAF) over the duration of the trial for interventions. Mean net change is the change from baseline to end point in the intervention group minus the change from baseline in the control group, or mean net change between baseline and end point for single-arm studies. Outcomes extracted for observational studies were a comparison of outcome measurements between dietary patterns.

#### **Quality assessment**

The quality of included studies was assessed independently and in duplicate using the Academy of Nutrition and Dietetics Quality Criteria Checklist (Table 3).<sup>80</sup> Four relevance questions and 10 quality questions were rated yes or no, ranging from clarity of research question, selection bias, randomization, dropout, blinding, clarity of intervention description, validity of measures, appropriateness of statistical analyses, and conclusions drawn and funding sources. A positive score was determined by "Yes" answers to questions 2, 3, 6, and 7, and at least 1 additional "Yes" on the other questions. If a "No" was the answer to 1 of questions 2, 3, 6, and 7 overall, and there were  $\geq 8$  "Yes" answers, the study was rated positive. If answers to 2, 3, 6, and 7 were "No," the study was rated as neutral. The study received a negative score if  $\geq 6$  of the 10 questions were responded to with "No."

# Data synthesis

A quantitative synthesis of the data was unable to be performed because of substantial diversity in methodology, dietary patterns, and measurements for outcomes of interest. As such, a narrative review was performed.

# Meta-bias(es)

To assess whether reporting bias was present in intervention studies, an investigation of whether each study's protocol had been published before commencement of the trial was undertaken. For all studies published after July 1, 2005, the Clinical Trial Register of the International Clinical Trials Registry Platform of the World Health Organization was searched and outcome reporting bias was assessed on the basis of whether selective reporting of outcomes were present.

#### RESULTS

Figure 3 presents the process and PRISMA flowchart for study selection. After deduplication, we identified 652 articles through the literature search. After reviewing titles and abstracts, 56 articles were relevant for fulltext review. Exclusion of full-text articles was based largely on the lack of examination of a dietary pattern. Sixteen articles were eligible and included for narrative synthesis.

Table 2<sup>17,81-95</sup> lists the characteristics of included studies. The majority of studies were undertaken in Greece (n = 5) and the United States (n = 3). Two studies were undertaken in South Korea and 1 each in Taiwan, India, Sweden, Iran, Spain, and Canada. Specific dietary patterns identified in the literature included "Mediterranean" dietary patterns, "vegetarian" dietary patterns, and "other heart healthy" dietary patterns (which included the Dietary Approaches to Stop Hypertension, or DASH, pattern; Living Heart dietary pattern; National Cholesterol Education Program dietary pattern; and a dietary pattern that replaced refined carbohydrates with whole grains and legumes and more vegetables). A posteriori dietary patterns were also reported and highlighted different patterns consumed across different population groups (namely in Greece, Sweden, and Iran). Data relating to primary and secondary outcomes were extracted from 7 randomized controlled trials (RCTs), 2 non-RCTs, 2 pre-post or single-arm studies, and 1 fixed-sequence intervention trial. The remaining 4 studies were cross-sectional.

In the 4 intervention studies examining Mediterranean dietary patterns, 2 showed significant reductions in PAF-induced aggregation of platelets in both healthy participants and people with type 2

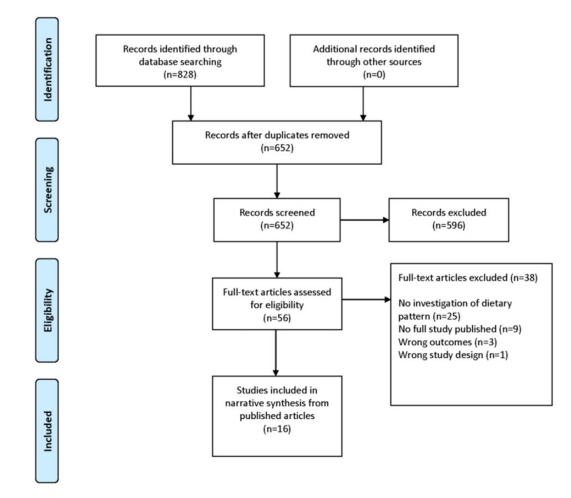


Figure 3 PRISMA flowchart of article selection.

diabetes, with the latter showing a much greater response.<sup>17,88</sup> A post hoc study of the Prevención con Dieta Mediterránea trial found a significant favorable change in Lp-PLA<sub>2</sub> activity levels in HDL after a 1-year Mediterranean dietary intervention supplemented with extra-virgin olive oil, when compared with a low-fat diet. However, no significant difference was seen in the Mediterranean diet group supplemented with nuts, when compared with a low-fat diet.<sup>81</sup> The other study was a fixed-sequence study that presented Lp-PLA<sub>2</sub> as percentage change only, which limited the usefulness of the data.<sup>91</sup> In that study, the small number of people whose HDL cholesterol was noted to have increased (n=6 compared with n=6 with reduced HDL), and there was a trend toward a favorable impact on Lp-PLA<sub>2</sub>; however, the results were not significant.<sup>91</sup>

Four studies examined vegetarian dietary patterns. One study was an RCT and compared similar Indian vegetarian diets that differed in the addition of either coconut or peanuts.<sup>83</sup> Results showed PAF reduced within the peanut group, but no between-group analysis was conducted.<sup>83</sup> In the single cross-sectional study in Taiwan,<sup>95</sup> Lp-PLA<sub>2</sub> activity was less favorable in omnivores. However, overall, both groups had low average Lp-PLA<sub>2</sub> levels, which could be due to Asian ethnicity.<sup>96</sup> In the 2 papers that reported pre-post single-arm studies, 1 reported significantly lower Lp-PLA<sub>2</sub> levels after 4 weeks of a raw, vegan dietary intervention.<sup>89</sup> The other reported a marginally significant increase in Lp-PLA<sub>2</sub> after 21 days of a largely vegetarian Pritikin dietary pattern.<sup>90</sup>

Heart-healthy dietary patterns were investigated in 5 studies, 4 of which were RCTs. Two of the RCTs focused on the replacement of refined grains with whole grains, increased vegetables, and addition of legumes in a South Korean population sample.<sup>84,85</sup> There were significant reductions in Lp-PLA<sub>2</sub> levels after a 12-week intervention. Another RCT evaluated a 3-week hearthealthy dietary pattern (the Living Heart Diet) combined with exercise and found significant reductions in Lp-PLA<sub>2</sub> compared with participants receiving usual care.<sup>86</sup> A pre-post study with a heart-healthy dietary intervention that was broadly similar to the Living Heart Diet found no significant difference in Lp-PLA<sub>2</sub> levels

Reference and study location Shankar (2017) RCT India <sup>83</sup>	Study design	Inclusion criteria			Distantion fistomication	Control	Outcomes (measurement method)
		ווורוחזוחוו רוורביים	Population mean ±SD or	Duration	Dietary pattern/intervention		
			(range)				mean ± SD or (range) <sup>a</sup>
India		Healthy adults	n = 58 (31 M, 27 F) Age:	90 d	n = 27 Vegetarian dietary pattern	n=31 Vegetarian dietary pattern	PAF pg/mL (ELISA) Vegetarian
			$23.8\pm4.8$ y Coconut group		with Coconut group: Balanced	with Peanut group: Balanced	with coconut group: Pre:
			weight: 59.8 $\pm$ 10.2 kg		vegetarian Yogic diet (based on	vegetarian Yogic diet $+$ 45 g	$186.88 \pm 383.11$ Post:
			Peanut group BMI:		grains, pulses, fruits, and vege-	peanuts $+$ 22 g/d peanut oil	$194.52 \pm 174.40; P = 0.947$
			$56.8\pm7.3~\mathrm{kg/m^2}$		tables) + 100 g/d fresh coconut		Vegetarian with peanut
							group: Pre: $375.25 \pm 705.03$
							Post: 139.45 $\pm$ 144.8;
							<b>P</b> = 0.05 Between-group dif-
							ference: $P = 0.224$ PON1 ng/
							<i>mL</i> Vegetarian with Coconut
							group: Pre: 2679.78 ± 878.8
							Post: 2755.82 ± 918.3;
							P= 0.67 Vegetarian with pea-
							nut group: Pre:
							2221.68 ± 647.7 <b>Post:</b>
							2773.59 ± 1145.7;
							<i>P</i> = 0.001 Between-group dif-
							ference: $P = 0.95 MPO nq/mL$
							Vegetarian with Coconut
							aroun: Pre: 657 92 + 599 22
							9.000
							P = 0.84 Vegetarian with
							Peanut group: Pre:
							648.57 ± 529.38 <b>Post:</b>
							924.26 ± 724.24; P = 0.006
							Between-group difference: P
							= 0.17
Kim et al RCT		Nonobese adults with	n=80 (M:F ratio: not	12 wk	n = 40  Whole-arain dietary pattern	n = 40 Usual diet (control) aroun:	Plasma Ln-PLA, activity (nmol/
		imnaired fasting	renorted) Are: 40_70 v		Whole-grain diet group: Refined	lisual Korean diet with refined	mil/min) (high-throughout ra-
(2010) 5			Michael and and a second and		viriore grant aret group: henrica		
South		glucose or newly	weight: not reported BMI:		rice replaced with 33% legumes,	rice	aiometric assay) Whole-grain
Korea <sup>84</sup>		diagnosed	not reported		33% barley, 33% wild rice $3 \times /d$		diet group: Pre: 28.0 $\pm$ 1.2
		diabetes			+6 servings of vegetables		Post: 25.7 ± 1.11; P > 0.05
					(180–420 g)		Usual diet group: Pre:
							$30.1 \pm 1.64$ Post: $30.3 \pm 1.61$ ;
							P > 0.05 Between-group dif-
							ference (change adjusted
							for baseline): $P < 0.001$
							Unstimulated PBMC Lp-PLA,
							activity (nmol/ml /min) Whole-
							drain diet droum: Pre-
							9.400 acc 9.446

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	Study design	Inclusion criteria	Population mean ±SD or	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method)
study location			(range)				mean $\pm$ SD or (range) <sup>a</sup>
							diet group: Pre: 2.00 $\pm$ 0.12
							Post: 2.28 ± 0.13; <i>P</i> < 0.01
							Between-group difference
							(change adjusted for base-
							<b>line):</b> $P < 0.001$ LDL particle
							size (nm) Whole-crain diat
							size (nini) winole-grain alet
							group: Pre: 24.4 $\pm$ 0.15 <b>Post:</b>
							24.6 ± 0.17; P < 0.001 Usual
							diet group: Pre: 24.1 $\pm$ 0.12
							Post 24.1 $\pm$ 0.13; $P > 0.05$
							Between-aroup difference
							(change adjusted for base-
							line): <i>P</i> = 0.001
Kim et al RCT		Adults with impaired	n = 99 (67 M. 32 F) Age. v:	12 wk I	n=50~Whole-arain dietary pattern	n = 49 Usual diet (control) group:	Plasma Lp-PLA, activity (nmol/
		fasting glucose.			Whole-grain diet group: Befined	Usual Korean diet with refined	ml /min) (hiah-throughout ra-
South		impaired alucose	56.3 + 1.2 Usual diet (con-		rice replaced with 33% legumes	rice	diometric assav) Whole-arain
V. 0000 85		intolorance or	++++++++++++++++++++++++++++++++++++++		Which with a state of the state	8	
NOIEd							
		newly diagnosed	not reported BMI (in lieu of		33% wild rice $3 \times / d + 6$ servings		Post: 27.8 ± 1.08; P < 0.01
		T2DM	weight): Whole-grain diet		of vegetables (180–420 g)		Usual diet group: Pre:
			group: 24.0 $\pm$ 0.38 kg/m <sup>2</sup>				$29.16 \pm 1.29$ Post:
			Usual diet (control):				$29.84 \pm 1.28; P > 0.05$
			$24.1 \pm 0.44 \text{ kg/m}^2$				Between-group difference
			ı				(change adjusted for base-
							line): P < 0.001 Unstimulated
							PBMC Lp-PLA <sub>2</sub> activity (nmol/
							<i>mL/min</i> ) Whole-arain diet
							drollin: Pre: 2 15 + 0 11 <b>Post:</b>
							1 86 ± 0 11: P < 0 001 Ileual
							diat around Drov 1 00 + 0 1 1
							Dest: 2.27 + 0.13: P < 0.01
							between-group difference
							(change adjusted for base-
							<b>line): P &lt; 0.001</b> <i>LDL particle</i>
							<i>size (nm)</i> Whole-grain diet
							group: Pre: 24.3 ± 0.12 <b>Post:</b>
							24.5 ± 0.14; P < 0.01 Usual
							diet aroun. Pre: 24 11 + 0 10
							Post. 24.01 + 0.14. P > 0.05
							(ulariye aujusted for base-
				24 wk			line): r = 0.048
							(continued)

Reference and study location	Study design	Inclusion criteria	Population mean ±SD or (range)	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method) mean $\pm$ SD or (range) <sup>a</sup>
			,				•
Wooten et al	RCT (5-arm drug trial) Data	Dyslipidemic, HIV-pos-	n = 107 (98 M, 9 F) Age:		n=22 Heart Healthy dietary pat-	n = 19 Usual care (control) group:	Lp-PLA <sub>2</sub> mass (ng/mL <sup>1</sup> )
(2013)	extracted for 2 arms only: (1)	itive adults treated	44.8 $\pm$ . 9 y Weight: Living		tern, Living Heart Diet group:	General advice on heart-healthy	mean $\pm$ SE (ELISA, PLAC test)
United	Living Heart Diet group (diet	with highly active	Heart Diet 81.6 $\pm$ 2.0 kg		Carbohvdrate. 50% energy: fat.	diet and exercise + 2 placebos.	Living Heart Diet group: Pre:
Statec <sup>86</sup>	(unitation on medication)	antiratroviral	lleual care (control)		30% anarciv (/ 7% SFA 15%	Darticipants diven booklet titled	387.0 + 17.0 <b>Deet</b> -
ומובס							
	and (2) usual care (control) only	therapy	$78.4\pm1.9~ m kg$		MUFA, 8% PUFA, minimal TFA),	Nutrition and Your Health	<b>323</b> ± <b>27.2;</b> P < 0.05 Usual
					cholesterol < 200 mg/d, fiber		care (control) group Pre:
					20-30  g/d + 2  placebos. Aerobic		415.1 ± 31.7 Post:
					and resistance exercise: 75–		$402.2 \pm 25.3; P > 0.05$
					90 min 3×/wk.		Between-aroup difference
							(adjusted for baseline). D
							CU.US NAIVIES (IIG/IIIL )
							<i>mean</i> ± <i>SE</i> Living Heart Diet
							group: Pre: 40.0 $\pm$ 3.2 Post:
							$55.0 \pm 11.3$ ; $P > 0.05$ Usual
							care (control) group: Pre:
							$42.4 \pm 5.9$ Post: 50.9 $\pm 10.4$ :
							P > 0.05 Between-group dif-
							ference (adjusted for base-
							line): <i>P</i> > 0.05
Rizos et al	RCT: only cross-sectional data	Adults with impaired	n = 151 (73 M, 78 F) Age: 60	12 wk	n = 151  DASH dietary pattern: all	N/A	Cross-sectional data extracted
(2011)	extracted Results extracted for	fasting plasma glu-	(46–70) y Weight: not		groups		Plasma Lp-PLA <sub>2</sub> activity (nmol/
Greece <sup>87</sup>	baseline data only (all 3 arms),	cose, mixed dysli-	reported BMI (in lieu of				mL/min) (TCA precipitation)
	after dietary intervention but	pidemia, and stage	weight): Group 1: 29 $\pm$ 4				Group 1 (RT): 57 $\pm$ 17 Group 2
	before randomization to drug	1 hypertension	kg/m <sup>2</sup> Group 2: 29 $\pm$ 5 kg/				(RI): 53 ± 11 Group 3 (RO):
	interventions	:	$m^{2}$ Group 3: 28 ± 4 kg/m <sup>2</sup>				$58 \pm 14$ Plasma Lp-PLA, mass
							(ng/ml) (ELISA, PLAC TEST)
							Group 1: 277 $\pm$ 40 Group 2:
							301 ± 20 Group 3: 304 ± 34
							Small dense LDL cholesterol
							(mg/dL) [mmol/L), median
							(range)] Group 1: 17 (2–69)
							[0.4 (0.1–1.8)] Group 2: 15 (7–
							44) [0.4 (0.2–1.1)] Groun 3: 17
							(2–78) [0.4 (0.1–2)] LUL Parti-
							cle size (A) Group 1: 261 $\pm$ 7
							Group 2: 262 ± 4 Group 3:
							$262 \pm 6$
Karantonis et al	Non-RCT	T2DM: managed with	n = 67 (35 M, 32 F) Age: 56	4 wk	Total n = 45 2 groups: Healthy:	Total $n = 22$ (T2DM: all) Usual diet	PAF EC <sub>50</sub> (PAF-induced platelet
(2005		diet or OHAS.	(26–74) v Weiaht: 77 ± 9 ka		n = 22: T2DM: $n = 23$ ]		aaareaation in PRP) Healthy
Greece <sup>88</sup>		Healthy age- and			Mediterranean-type dietary nat-		droun: Pre: 1 45 + 1 47 Post:
		weight-matched			tern: based on rast-rood meals		2./0 ± 2.39; P = 0.023
		adults			pretested for ability to reduce		T2DM aroup: Pre: 1.02 ± 1.38

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Reference and study location	Study design	Inclusion criteria	Population mean ±SD or (range)	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method) mean $\pm$ SD or (range) <sup>a</sup>
					PAF-induced aggregation		Post: 2.40 ± 4.65; <i>P</i> = 0.019
					in vitro (TPL)		Usual/control (T2DM) group:
							Pre: 0.774 $\pm$ 0.522 Post:
							$0.831 \pm 0.5; P = 0.285$
t al	Single-arm trial	Overweight or obese	n = 22 (22 M) Age : 62.8 (46–	21 d	n=22 Vegetarian dietary pattern	N/A	PAF-AH activity (nmol PAF/min/
(2006)		adult males	76) y Weight: 103.4 $\pm$ 22.9		Low-fat, Pritikin diet $\geq$ 5 serv-		mg protein) (solid-phase chro-
USA <sup>90</sup>			kg		ings/d whole grains, $\geq$ 4 serv-		matography with liquid scintil-
					ings/d vegetables $\geq$ 3 servings/d		<i>lation)</i> Pre: 23.4 $\pm$ 0.6 <b>Post:</b>
					fruit. Protein from plant sources,		24.6 $\pm$ 0.6; P = 0.05 $PON1$
					nonfat dairy $\leq$ 2 servings/d;		activity per mg/HDL Pre:
					fish/fowl 85–140 g/wk. Minimal		669.2 ± 95.6 Post:
					SFA and trans FA intake; no		$684.8 \pm 99.7; P > 0.05$
					added fats, sugars $+$ 45–60 min		
مدالب بفر المحمالفر سورط					walking/d		
Ubservational studies Hlebowicz et al Prosp	lies Prospective cohort study	Adult men and	n = 4999 (2040 M; 2959 F)	N/A	n = 4999 A posteriori dietary pat-	N/A	General linear model (controlled
		women No diagno-			terns identified by cluster analy-		for age, total energy, season,
Sweden <sup>94</sup>		sis of diabetes (IFG	v Weight: not reported		sis Six dietary patterns 1. Many		% body fat. WHR) LD-PLA,
		eligible) or previ-	-		foods and drinks 2. Fiber-rich		mass (ng/mL <sup>1</sup> ) (ELISA, PLAC
		ous history of CVD			bread 15% of energy from fiber-		test) Many foods and drinks
					rich bread 3. Low-fat and high-		pattern (n = 1399): <b>Male:</b>
					fiber foods 10.5% of total en-		287.39 <u>+</u> 3.76 Female:
					ergy from fruit, 8% from low-fat		<b>258.72</b> ± <b>2.65</b> Fiber-rich
					milk, both high-fat and low-fat		bread pattern (n = 460):
					meats and sweets 4. White		Male: 286.51 ± 5.48
					bread 16% of total energy from		Female: 257.15 ± 5.17 Low-
					white bread, other major energy		fat and high-fiber foods pat-
					sources were low-fat margarine,		tern (n = 755): <b>Male:</b>
					both high-fat and low-fat meats		284.55 ± 6.97* Female:
					and sweets 5. Milk-fat pattern		<b>250.64</b> ± <b>3.26</b> * White-bread
					12% of total energy from but-		pattern (n = 713): Male:
					ter/rapeseed oil spread, other		291.74 ± 4.22 Female:
					major energy sources included		<b>263.62</b> ± <b>4.40</b> Milk-fat pat-
					cheese, whole milk, + some		tern (n = 638): <b>Male:</b>
					white bread and sweets 6.		308.03 ± 4.84** Female:
					Sweets and cakes pattern 18%		<b>269.25</b> ± <b>4.23</b> ** Sweets and
					of total energy from sugar,		cakes pattern (n = 1034):
					sweets, jam; other major energy		Male: 296.33 ± 4.17
					sources were cakes, biscuits, and		Female: 265.42 ± 3.19
					soft drinks		Male: $P = .009$ ; Female:
							$\mathbf{P} = 0.004 \text{ L}p\text{-}PLA_2$ activity
							(na/mL <sup>1</sup> ) (hiah-throuahput

# (continued)

study location		Population mean ±5D or	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method)
		(range)				mean $\pm$ SD or (range) <sup>a</sup>
						radiom etric assay) Many
						foods and drinks pattern
						(n = 1399): Male: 49.17 $\pm$ 0.61
						<b>Female: 41.59</b> ± 0.42* Fiber-
						rich bread pattern (n $=$ 460):
						Male: 50.70 $\pm$ 0.89 (lowest as-
						sociation) Female:
						$42.98 \pm 0.82$ Low-fat and
						high-fiber foods pattern
						(n = 755): Male: 47.58 ± 1.13
						(highest association) Female:
						$42.01\pm0.52$ White-bread
						pattern (n $=$ 713): Male:
						49.89 ± 0.68 <b>Female:</b>
						<b>44.06</b> $\pm$ <b>0.70</b> (highest associ-
						ation) Milk-fat pattern
						(n = 638): Male: 50.09 $\pm$ 0.78
						Female: 43.27 ± 0.67 Sweets
						and cakes pattern (n $=$ 1034):
						Male: 49.93 ± 0.67 <b>Female:</b>
						<b>43.40</b> ± <b>0.51</b> Male: <i>P</i> = .291
						Female: $P = 0.007$
Chen et al Cross-sectional	Healthy, adult, non-	n = 363 (363 F) Age:	N/A	n = 173 Vegetarian dietary pattern	n = 190  Omnivore dietary pattern	Lp-PLA <sub>2</sub> activity 10 <sup>-3</sup> µmol/min/
(2011)	smoking women	$51.9 \pm 9.9$ y Weight: not		Lacto-ovo vegetarian		mL (PAF acetylhydrolase color-
Taiwan <sup>95</sup>		reported BMI (in lieu of				<i>imetric assay</i> ) Vegetarian:
		weight): Omnivores:				18.32 <u>+</u> 7.19 Omnivore:
		$23.28 \pm 3.47 \text{ ka/m}^2$				20.22 ± 8.13 Between-group
		Veaetarians: 22.87 ± 2.94				difference: $P < 0.05$
		-9-				Initiation linear representation
		Kg/III				
						Vegetarian: $\beta = -0.19$
						(-3.63, 0.016); <i>P</i> < 0.05
						Multivariate regression (age
						and BMI) Vegetarian: $\beta$ =
						-1.79 (-3.58, -0.01):
						P < 0.05
Intervention studies						
Hernaez et al RCT	T2DM or $\geq$ 3 cardio-	n = 358 (131 M, 227 F)	1 y	Total $n = 239$	Total $n = 119$	PAF-AH activity in HDLs (PAF
(2020)	vascular risk factors	Age : 66.8 $\pm$ 5.8 y		2 groups:	Low-fat diet	acetylhydrolase colorimetric
Spain <sup>81</sup>	(cholesterol, hyper-	Weight: not reported		Mediterranean diet supplemented		assay) (1-y change):
	tension, BMI,	BMI: mean not reported		with EVOO: $n = 120$ ;		Mean change (95%Cl)
	smoking, family			Mediterranean diet supple-		Mediterranean diet with EVOO vs
	history			mented with nuts: $n = 119$		control:

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Reference and study location	Study design	Inclusion criteria	Population mean ±5D or (range)	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method) mean ± SD or (range) <sup>a</sup>
							<b>7.48% (0.17–14.8)</b> Mediterranean diet with nuts vs control: 3.39% ( – 3.64 to 10.4)
Makariou et al (2019) Greece <sup>82</sup>	RCT Results extracted for single-arm control group only (diet + no supplement)	Adults with metabolic syndrome	n = 50 (25 M, 25 F) Age: 53 (37–67) y Weight: 89.0 ± 13.4 kg	е ж	n = 25 Hear Healthy Dietary Pattern NCEP ATP III guidelines Fat 25-35% energy (< 7% 5FA, re- duced TFA), dietary cholesterol < 200 mg/d. Most dietary fat unsaturated; simple sugars limited	N/N	Heart-thealthy distary pattern Heart-thealthy distary pattern $Lp-PLA_2$ activity (mmol/mL/min) (TCA precipitation) Pre: 57.4 $\pm$ 13.3 Post: 52.7 $\pm$ 12.4; $P > 0.05$ sdLDL cholesterol mg/dL Pre: 7 (0-22) Post: 5 (2-25); $P > 0.05$ sdLDL proportion, % Pre: 38 $\pm$ 2.3; $P > 0.05$ Mean LDL size (nm) Pre: 266.5 $\pm$ 3.5; $P > 0.05$
Antonopoulou et al (2006) Greece <sup>17</sup>	Non-RCT	Type 2 diabetes: man- aged with diet or OHAs. Healthy age- and weight-matched adults	n = 69 (37 M, 32 F) Age: 53 (26-70) y Weight: 77 ± 9 kg	4 WK	Total $n = 46$ 2 groups: Healthy: $n = 22$ ; T2DM: $n = 24$ ] Mediterranean-type dietary pattern: Based on catering company-sup- plied meals pretested for ability to reduce PAF aggregation in vitro (TL)	Total n = 23 (T2DM: all) Usual diet	PAF EC <sub>50</sub> (PAF-induced platelet aggregation in PRP) Healthy group: Pre: $1,4 \pm 1,4$ Post: $2,70 \pm 2.6$ ; $P = 0.023$ T2DM group: Pre: $0.76 \pm 0.5$ Post: $4,2 \pm 1.2$ ; $P < 0.001$ Baseline significantly different be- tween groups Usual/control (T2DM) group: Pre: $0.77 \pm 0.52$ Post: $0.83 \pm 0.5$ ; $P = 0.285$
Najjar et al (2018) United States <sup>89</sup>	Single-arm trial	Adults with hypertension and dyslipidemia: dyslipidemia: SBP $\geq$ 140 mmHg or DBP $\geq$ 90 mmHg, LDL-C $\geq$ 100 mg/dL and BMI $\geq$ 255 kg/m <sup>2</sup> .	n = 31 (10 M, 21 F) Age: 53.4 (32–69) y Weight: 108.1 ± 5.1 kg	4 wk	n = 31 Vegetarian dietary pattern (vegan, raw) Vegan, raw plant-based diet. <i>raw</i> <i>fruits, vegetables, avocado, seeds,</i> <i>and plant foods dehydrated to</i> <i>temperatures ≤ 160° F ad libitum.</i> <i>Cooked foods, animal products,</i> <i>free oils, soda, alcohol, and coffee</i> <i>were excluded.</i>	N/A	<ul> <li>Lp-PLA<sub>2</sub> mass (ng/mL) (not reported)</li> <li>Vegan raw plant-based diet:</li> <li>Pes: 252.3 ± 136.3</li> <li>Post: 210.7 ± 119.1; P = 0.001</li> <li>MPO (pmol/L)</li> <li>Pre: 124.1 ± 58.1</li> <li>Post: 104.5 ± 536, P = 0.056</li> <li>sdLDL cholesterol mg/dL</li> <li>Pre: 33.7 ± 11.5</li> <li>Post: 33.7 ± 11.5</li> </ul>

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Reference and study location	Study design	Inclusion criteria	Population mean ±SD or (range)	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method) mean $\pm$ SD or (range) <sup>a</sup>
Richard et al (2014) Canada <sup>91</sup>	Fixed-sequence intervention	Nonsmoking male adults with meta- bolic syndrome No CHD or diabetes; not taking lipid- lowering or antihy- pertensive	n = 26 (26 M) Age: 49.4 (24–62) y Weight: 98.3 ± 17.6 kg	10 wk	n = 26 Mediterranean dietary pattern 5-wk controlled feeding interven- tion: high in whole grains, legumes, fruits, vegetables, fish, olive oil, ruts, and moderate amoutt of red wine amoutt of red wine	n = 26 Standard North American diet— the intervention diet followed a 5-wk run-in, which served as the control	PAF-AH HDL protein (fold change) (mass spectrometry iTRAQ) Med diet vs control = 1.10; P = 0.845 error factor = 5.93 (an error factor value > 2 indicates the ratios
Seyedi et al (2020) Iran <sup>92</sup>	Cross-sectional	medication Adult men and women ≥5 of: TC >200 mg/ dL, LDL C >100 mg/dL, HDL C <40 mg/dL (M), <50 mg/dL (F), waist circ. = >102 cm (M), >88 cm (F), SBP >102 cm (M), >88 cm (F), SBP >102 cm Hg, anti- hypertensive medi- cation, age ≥45 y (M), >55 v (F).	n = 470 (114 M, 356 F) Age: 40-70 y Weight: not reported	N/A	<ul> <li>n = 470</li> <li>A posteriori dietary pattern identified by factor analysis. Three dietary patterns calculated:</li> <li>1. Healthy (reference pattern): high in fresh and dried fruits, olives, high-and low-fat dairy products, poultry and fish, liquid oils, and canned products</li> <li>2. Semi-Mediterranean: characterized by legumes, potatores, eggs, red meats, tea, and coffee.</li> <li>3. Western:</li> </ul>	K N	vary greatly from peptide to peptide) $(\mu P. RA_3 mass ng/mL (ELSA))$ Univariate linear regression Western: $\beta = 0.35 (0.11, 0.78); P = 0.026$ Semi-Mediterranean: $\beta = 0.35 (0.11, 0.78); P = 0.026$ Semi-Mediterranean: $\beta = -0.12 (-3.52, -0.16);$ P = 0.043 Mutivariate linear regression (age, BMI, activity, El, FBG, hormone therapy, lipid-lowering drugs) Western: $\beta = 1.32 (1.05, 1.64); P = 0.035$ Semi-Mediterranean $\beta = -0.01 (-0.16, 0.43); P = 0.75$
Detopoulou et al (2015 Greece <sup>03</sup>	Cross-sectional	smoker Healthy adults No history of CVD or inflammatory dis- ease, no current respiratory infec- tion, dental prob- lems, renal/hepatic abnormalities. Men were age- and BMI-matched to women.	n = 106 (48 M, 58 F) Age : 44 (31-57) y Weight: not reported BMI (in lieu of weight): 27.5 kg/m <sup>2</sup>	NA	fast foods, salty snacks, mayon- naise, and organ meats Mediterranean Dietary Pattern (and 2 miscellaneous other patterns): 1. A priori MedDietScore (as devel- oped by Panagiotakos et al, 2006): based on nonrefined cereal, fruits, vegetables, potatoes, legumes, olive oil, fish, red meat, poultry, full-fat dairy products, and alcohol). 2. Calculation of dietary antioxidant capacity 3. Six a posteriori dietary patterns identified by principal compo- nent analysis 1: Fruits, nuts, and herbal drinks	None	Total PAF (fmol/mL), median (lower-upper quartile) (PAF-in- duced platelet aggregation to- ward washed rabbit platelets) Male: 82 (29–372) Female: 152 (43–944) Total: 119 (34–578) MedDiet5core: Men only (n = 48); Adjusted for age, sex, El/BMR Bound PAF r = -0.36, $P = 0.08Total PAFr = -0.30$ , $P > 0.05Dietary antioxidant capacity: ad-justed for age, sex, El/BMRTotal PAF (pmol/mL)$

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Reference and St	Study design	Inclusion criteria	Population mean ±SD or	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method)
study location			(range)				mean $\pm$ SD or (range) <sup>a</sup>
				2:	2: Legumes, vegetables, poultry		DAC FRAP: $r = -0.197$ ; $P = 0.06$
					and fish		DAC-TRAP: $r = -0.211$ ; $P = 0.04$
				3:	3: Low consumption of low-fat		DAC TEAC: $r = -0.200$ ; $P = 0.05$
					dairy, high consumption of full-		Lyso-PAF-AT (nmol/min/mg)
					fat dairy, cheeses, alcohol, and		DAC FRAP: $r = -0.200$ ; $P = 0.05$
					red meat		DAC-TRAP: $r = -0.171$ ; $P = 0.1$
				4:	4: Coffee and low intake of whole-		DAC TEAC: $r = -0.146$ ; $P = 0.1$
					wheat products		Lp-PLA <sub>2</sub> (nmol/min/mL) (TCA
				5:	5: Refined cereals and full-fat dairy,		precipitation)
					cheeses		DAC FRAP $r = 0.090$ ; $P = 0.30$
				9:	6: Whole-wheat products and olive		DAC TRAP $r = 0.119$ ; $P = 0.20$
					oil		DAC TEAC $r = 0.110$ ; $P = 0.30$
							Free PAF, bound PAF, PAF-CPT, and
							PAF-AH: all results not significant.
							A posteriori dietary patterns:
							Linear regression adjusted for age,
							sex, El/BMR, and other dietary
							patterns
							Free PAF pmol/mL
							Legumes, vegetables, poultry, and
							fish dietary pattern:
							$-0.157 \pm 0.087; P = 0.07$
							Total PAF pmol/mL
							Coffee and low intake of whole-
							wheat products dietary pattern:
							$-0.147 \pm 0.08; P = 0.06$
							Lyso-PAF-AT (nmol/min/mg)
							Fruits, nuts, herbal drinks:
							$-1202 \pm 652; P = 0.06$
							Whole-wheat products, olive oil di-
							etary pattern:
							$-1273 \pm 571; P = 0.02$
							Cox proportional hazards regres-
							sion (adjusted for age, total en-
							ergy, season, % body fat, WHR,
							and smoking)
							Tertile 1: lowest adherence; tertile 3:
							highest adherence
							Lp-PLA <sub>2</sub> mass (ng/mL <sup>1</sup> )
							Female:
							Low-fat and high-fiber foods
							pattern:

(continued)

Reference and	Study design	Inclusion criteria	Population mean ±SD or	Duration	Dietary pattern/intervention	Control	Outcomes (measurement method)
study location			(range)				mean $\pm$ SD or (range) <sup>a</sup>
							Tertile 2: OR, 0.89 (0.71, 1.12)
							Tertile 3: OR, 0.69 (0.54, 0.87)
							P = 0.002
							Sweets and cakes pattern:
							Tertile 2: OR, 1.20 (0.96, 1.50)
							Tertile 3: OR, 1.29 (1.02, 1.62)
							P = 0.030
							No significance when those with
							past change in diet were ex-
							cluded ( $P = 0.098$ and $P = 0.149$ ,
							respectively)
							Data for other patterns not
							reported
							Lp-PLA <sub>2</sub> activity (ng/mL <sup>1</sup> )
							Male:
							Low-fat and high-fiber foods
							pattern:
							Tertile 2: OR, 0.92 (0.61, 1.38)
							Tertile 3: OR, 0.62 (0.40, 0.96)
							P = 0.036
							No significance when those with
							past change in diet were ex-
							cluded: $P = 0.352$
							Milk-fat pattern
							Tertile 2: OR, 1.17 (0.85, 1.62)
							Tertile 3: OR, 1.50 (1.10, 2.05)
							P = 0.011
							P = 0.009 when those with past
							change in diet were excluded
							Data for other patterns not
							renorted

MUFA, monounsaturated fatty acid; N/A, not applicable; OHA, oral hypoglycemic agent; OR, odds ratio; PAF, platelet activating factor; PBMC, peripheral blood mononuclear cells; PRP, platelet-rich plasma; PON1, serum paraoxonase and arylesterase 1; PUFA, polyunsaturated fatty acid; RCT, randomized controlled trial; SBP, systolic blood pressure; sdLDL, small dense low-density lipoprotein; SE, standard eror; SF, saturated fat; SFA, saturated fatty acids; T2DM, type 2 diabetes mellitus; TC, total Abbreviations: AH, acetylhydrolase; BMI, body mass index; BMR, basal metabolic rate; CHD, coronary heart disease; circ., circumference; CVD, cardiovascular disease; DAC, dietary antioxidant capacity; DASH, Dietary Approach to Stop Hypertension; DBP, diastolic blood pressure; EC<sub>50</sub>, half-maximal effective concentration; ELISA, enzyme-linked immunosorbent assay; EVOO, extra virgin olive oil; F, female; FA, fatty acid; FBG, fasting blood glucose; FRAP, ferric-reducing antioxidant power; HDL, high-density lipoprotein; IFG, impaired fasting glucose; iTRAQ, isobaric tags for relative and absolute quantitation; LDL-C, low-density lipoprotein cholesterol; Lp-PLA<sub>2</sub>, lipoprotein-associated phospholipase A2; M, male; MPO, myeloperoxidase; cholesterol; TCA, trichloroacetic acid; TEAC, trolox-equivalent antioxidant power; TFA, trans fatty acids; TRAP, total radical-trapping antioxidant parameters; WHR, waist to hip ratio.

<sup>3</sup>bold indicates statistically significant results P  $\leq$  0.05. For some observational studies, only statistically significant results (or results approaching significance) are included, for brevity.

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after 3 months.<sup>82</sup> In another RCT in which only crosssectional data were extracted, Lp-PLA<sub>2</sub> activity was reported after a 12-week Dietary Approaches to Stop Hypertension diet run-in period before randomization.<sup>87</sup>

Three cohort studies examined posteriori dietary patterns. One study in Sweden used cluster analysis to identify 6 novel dietary patterns, and the authors reported somewhat inconsistent findings across male and female participants.<sup>94</sup> However, across both sexes, the low-fat and high-fiber dietary pattern (10.5% of total energy derived from fruit, 8% energy from low-fat milk, both high-fat and low-fat meats, and sweets) was associated with lower Lp-PLA<sub>2</sub> levels, whereas the milk-fat pattern (12% of total energy derived from a butter/rapeseed oil spread and other major energy sources that included cheese, whole milk, and, to a lesser extent, white bread and sweets) was associated with higher Lp-PLA<sub>2</sub> levels.<sup>94</sup> A second study in Greece also identified 6 unique dietary patterns and found a pattern rich in whole-wheat products with olive oil was inversely correlated with levels of lyso-PAF acetyltransferase (an enzyme related to PAF metabolism).<sup>93</sup> In the same study, a high dietary antioxidant capacity score (but not a Mediterranean diet score) was inversely associated with total PAF after adjustment for confounders.<sup>93</sup> The third study identified 3 unique dietary patterns: (1)a healthy dietary pattern (ie, high in fruits, dried fruit, olives, high- and low-fat dairy products, poultry and fish, liquid oils, and canned products), (2) semi-Mediterranean dietary pattern (ie, legumes, potatoes, eggs, red meats, tea, and coffee), and (3) a Western dietary pattern (dominated by carbonated drinks, fast foods, salty snacks, mayonnaise, and organ meats).<sup>92</sup> Compared with the healthy dietary pattern, the Western dietary pattern was associated with less favorable Lp-PLA<sub>2</sub> levels. After accounting for confounders, the semi-Mediterranean dietary pattern showed no effect on Lp-PLA<sub>2</sub> with the healthy dietary pattern as the referent.

Four novel biomarkers were identified in the literature as secondary outcomes for this review: serum paraoxonase and arylesterase 1 (PON1), myeloperoxidase (MPO), RANTES (chemokine ligand 5; regulated on activation, normal T-cell expressed and secreted), and LDL particle size. PON1 is a cardioprotective enzyme that prevents the accumulation of oxidized LDL and promotes cholesterol efflux out of macrophages.<sup>97</sup> MPO is an enzyme linked to inflammation and oxidative stress and has been shown to be involved in all stages of atherosclerosis.<sup>98</sup> RANTES is a pro-inflammatory cytokine that induces leukocyte activation and migration and is associated with a wide range of inflammatory disorders.<sup>99</sup> LDL particle size can be a marker used in the prediction of CVD. Small dense LDL particles are a distinct LDL subclass that is more pro-atherogenic than large LDL particles because they have a decreased affinity for the LDL receptor, resulting in longer circulation time; enter the arterial wall more easily; are more prone to entrapment in the arterial wall; and are more susceptible to oxidation.<sup>100</sup>

A vegetarian diet supplemented with peanuts (but not the same diet supplemented with coconut instead of peanuts) resulted in a significant increase in PON1.<sup>83</sup> Similarly, MPO was significantly increased in the peanuts-supplemented group but not the coconut group.<sup>83</sup> The largely vegetarian Pritikin dietary pattern showed no effect on PON1 levels.<sup>90</sup>

Similarly, a raw vegan dietary pattern intervention significantly lowered small dense LDL particles and decreased levels of MPO (P = 0.056).<sup>89</sup> A heart-healthy intervention resulted in no significant difference in RANTES in either the usual-care or intervention groups.<sup>86</sup> LDL particle size was significantly increased in the whole-grain dietary pattern interventions compared with a refined-grains dietary pattern.<sup>84,85</sup>

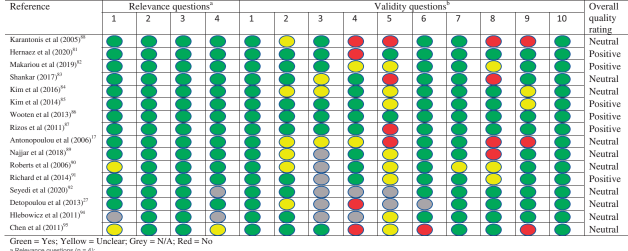
Risk-of-bias assessment identified 6 positive, 10 neutral, and 0 negative articles (Table 3). Studies that rated lower on the scale did so mostly because of inadequate description of follow-up methods and handling of withdrawals and methods of blinding. There were no discrepancies in outcome reporting when study reports were checked against the Clinical Trial Register of the International Clinical Trials Registry Platform of the World Health Organization.

## DISCUSSION

In this systematic review, we investigated the association between overall dietary patterns and their effect on PAF and Lp-PLA<sub>2</sub> as novel biomarkers of inflammation. There was a small number of published dietary studies reporting these biomarkers. Thirteen of the 16 included studies reported Lp-PLA<sub>2</sub> and only 4 reported PAF, with 1 study reporting on both markers. The paucity of research in this area is likely due to the novelty of the markers, in addition to the difficulty in measuring them and a lack of an established reference range for PAF and Lp-PLA<sub>2</sub> activity in a normal, healthy population.

However, a key finding from this review is that a range of established dietary patterns broadly consistent with country-specific dietary guidelines around the world show promise in producing favorable changes in these novel biomarkers. These included Mediterranean dietary patterns, vegetarian dietary patterns, and other heart-healthy dietary patterns. Conversely, dietary patterns including foods that were more highly processed

#### Table 3 Risk-of-bias assessment



a Relevance questions (n = 4): 1. Would implementing the studied intervention or procedure (if found successful) result in improved outcomes for the patients/clients/population group?

Did the authors study an outcome (dependent variable) or topic that the patients/clients/population group would care about?
 Is the focus of the intervention or procedure (independent variable) or topic of study a common issue of concern to dietetics practice?

4. Is the intervention or procedure feasible?

4. Is the intervention of procedure reasone :
b Validity questions (n = 10);
1. Was the research question clearly stated?
2. Was the selection of study subjects/patients free from bias?

Were study groups comparable? Was method of handling withdrawals described?

5. Was blinding used to prevent introduction of bias? 6. Were intervention/therapeutic regimens/exposure factor or procedure and any comparison(s) described in detail? Were intervening factors described?

Were outcomes clearly defined and the measurements valid and reliable?
 Was the statistical analysis appropriate for the study design and type of outcome indicators?
 Are conclusions supported by results with biases and limitations taken into consideration?
 Is bias due to study's funding or sponsorship unlikely?

and reflective of Western diets were associated with unfavorable outcomes.

The finding that Mediterranean dietary patterns were associated with favorable changes in levels of both PAF and Lp-PLA<sub>2</sub> post intervention is unsurprising. The Mediterranean diet was associated with reduced risk of CVD, including a reduction in events and deaths in a recent systematic review, although the effect size was small and the quality of evidence low to moderate.<sup>101</sup> A previous systematic review that investigated the Mediterranean diet or its components and PAF and Lp-PLA<sub>2</sub> found a range of foods to have favorable effects; the authors concluded that dietary patterns that emphasize cereals, legumes, vegetables, fish, and wine were worthy of additional investigation.<sup>30</sup> This study also noted that research was lacking on olive oil (the most characteristic component of Mediterranean diets). Although not specific to these novel biomarkers, another systematic review found that a Mediterranean dietary pattern was associated with lower levels of other markers of inflammation and improved endothelial function.<sup>102</sup> A Mediterranean diet intervention also significantly improved dietary inflammatory index scores (a measure of potential of diet to affect established inflammatory cytokines) compared with a low-fat diet in people with coronary heart disease.<sup>103</sup>

People with cardiometabolic conditions or risk factors may have greater responses to dietary intervention. Results from 2 studies we included in the present review suggested that Mediterranean dietary patterns may have greater favorable effects on PAF-induced platelet activity in patients with type 2 diabetes who are treated with both medication and diet, compared with healthy control study participants.<sup>17,88</sup> It is possible that this was due to lower platelet resistance to PAF-induced platelet aggregation in participants with type 2 diabetes at baseline, compared with healthy participants, which provides greater scope for improvement because of their naturally higher levels of platelet hyperactivity resulting in increased activation and aggregation.<sup>104</sup>

Furthermore, the results of the present study demonstrated that vegetarian dietary patterns were associated with more favorable changes in levels of PAF and Lp-PLA<sub>2</sub>. This is consistent with wider evidence supporting cardiovascular benefits of minimally processed plant-based diets, of which vegetarian dietary patterns are a subset.<sup>105</sup> Vegetarian diets emphasizing foods low in dietary fat may not confer the same benefits, because they are lower in fats that contain anti-inflammatory properties such as bioactive polar lipids (ie, phospholipids, sphingolipids, glycolipids) found in olive and seed oil, and higher-fat dairy products.<sup>20</sup> For example, in the Roberts study,<sup>90</sup> participants consumed non-fat milk that contained half the levels of PAF-inhibiting polar lipids than did whole milk.<sup>106</sup> Other research has highlighted potential benefits of full-fat dairy

consumption, due to a greater bioavailability of highvalue nutrients such as vitamin D and other antiinflammatory microconstituents.<sup>107,108</sup>

Within the current review, vegetarian diets with and without dairy and/or eggs were associated with favorable outcomes. One observational study found lower levels of Lp-PLA<sub>2</sub> in groups following a lacto-ovo vegetarian dietary pattern compared with groups who were omnivores; however, the former group had higher levels of high-sensitivity C-reactive protein than did the omnivore group.<sup>95</sup> These results are in contrast to those of a recent systematic review and meta-analysis that found vegetarian diets are associated with significantly lower levels of high-sensitivity C-reactive protein compared with nonvegetarian diets.<sup>109</sup> The researchers noted Taiwanese vegetarians consume fewer fresh vegetables, which they cook in oil, than do Western vegetarians, and they consume many deep-fried and refined soybean and grain products, which might contribute to higher high-sensitivity C-reactive protein levels.

The other heart-healthy dietary patterns associated with favorable effects on inflammation in this review are broadly similar to country-specific dietary guidelines across the United States, the United Kingdom, and Australia.<sup>110-112</sup> These guidelines advocate higher intakes of vegetables and fruits, moderate dairy consumption (albeit favoring reduced- or lower-fat options), plant-based oils, and unprocessed protein sources such as fish, lean meat, and legumes. A randomized dietary intervention study in healthy men and women compared a diet consistent with UK dietary guidelines with a representative UK diet and demonstrated a significant reduction in C-reactive protein levels after 12 weeks. This suggests that inflammation is positively affected when dietary guidelines are followed,<sup>113</sup> possibly via increased food sources of polyphenols,<sup>114</sup> known to be PAF inhibitors.<sup>63</sup> Research has shown an inverse association between Lp-PLA<sub>2</sub> and retinol and carotene, markers for provitamin A fruit and vegetable intake, in patients with incident CVD.115 Higher intake of fruit and vegetables led to a reduction in levels of inflammatory biomarkers in a recent systematic review and meta-analysis.<sup>116</sup>

We found that a Western dietary pattern is associated with higher levels of inflammation. This is not unexpected, because Western dietary patterns are associated with increased risk of coronary heart disease in both men and women,<sup>117,118</sup> and given the known link between inflammation and heart disease. A recent review found that Western dietary patterns are associated with increased levels of the blood inflammatory biomarkers high-sensitivity C-reactive protein, leptin, and IL-6.<sup>119</sup>

Very few secondary outcomes were identified in this review; however, key markers appear to be PON1, MPO, and LDL particle size. Results for these outcomes were mixed. LDL particle size appears to be an important predictor of cardiovascular events and small dense LDL particles are more pro-atherogenic than large LDL particles.<sup>100,120</sup> Levels of Lp-PLA<sub>2</sub> in small dense LDL have been reported to be 5 to 10 times higher than in normal-size LDL.<sup>121</sup> Of the 3 secondary outcomes, PON1 may be a useful addition to future studies investigating PAF and Lp-PLA<sub>2</sub>, given its presence within HDL and protective action against LDL oxidation.

Weight change may be a mediator of inflammatory biomarkers. Authors of a recent review (which did not include the novel biomarkers investigated in the present review) found no significant effect on markers of subclinical inflammation when examining whole foods and dietary patterns in weight-stable individuals with a high body mass index.<sup>122</sup> The review authors concluded that weight loss may be a key factor in dietary interventions that reduce inflammation. In the present review, there was no change in mean weight from baseline in 7 of 10 interventions, but there were improvements in inflammation after the interventions. Three studies noted significant weight loss, but inflammatory outcomes were inconsistent. One study<sup>89</sup> showed a weight loss of >6% of body weight after a 4-week intervention, with concomitant reductions in levels of novel inflammatory biomarkers. In contrast, the other 2 studies showed no or a worsening effect: one study<sup>87</sup> reported a small reduction in weight with no change in Lp-PLA<sub>2</sub> from baseline; the other study<sup>90</sup> reported a 3% reduction in body weight, but Lp-PLA<sub>2</sub> level actually increased after the intervention.

To our knowledge, this is the first systematic review to explore the association between dietary patterns, beyond the Mediterranean Diet, and the novel biomarkers PAF and Lp-PLA<sub>2</sub>. Strengths of our study include a strong methodology and use of the PRISMA guidelines. A comprehensive literature search was performed using 4 databases. Screening of title and abstracts and full-text review for inclusion criteria were performed in duplicate. Data extraction was independently reviewed for accuracy and quality assessment was performed.

This review was comprehensive and systematic; however, the analysis is limited by the small number of studies adhering to the inclusion criteria assessing dietary patterns and these novel biomarkers. The sheer novelty of the markers of interest are another limitation, because measurement methods are varied and no consensus of cutoff points have been derived for either PAF or Lp-PLA<sub>2</sub> activity, making it difficult to interpret the results reported in the studies. Other limitations of this study include the wide diversity of groups reported in the studies, which makes it difficult to draw comparisons, and the inclusion of cross-sectional studies that encompass a high risk of bias and lower level of study quality when compared with RCTs. The number of studies examining PAF was very limited, suggesting this is a gap in the literature. Large-scale intervention studies are needed to gain a better understanding of how diet affects this novel biomarker. Because little is known about the normal concentrations of both biomarkers in healthy populations, priority for research should be placed on establishing reference values to determine the clinical utility of these biomarkers.

#### CONCLUSION

There is limited evidence and considerable diversity in existing studies investigating dietary patterns and the novel inflammatory markers PAF and Lp-PLA<sub>2</sub>. A range of well-established dietary patterns has potential to improve these novel markers, including Mediterranean, vegetarian, and other heart-healthy dietary patterns. Conversely, Western dietary patterns are associated with higher levels of inflammation, as measured by these markers. More, well-designed studies are needed to confirm these findings and identify other dietary patterns that could positively affect inflammation.

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Declaration of interest. The authors declare no conflict of interest.

# **Supporting Information**

The following Supporting Information is available through the online version of this article at the publisher's website.

*Table S1* Search terms used in the PubMed, CINAHL, Embase, and Cochrane databases

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#### REFERENCES

- Frostegard J. Immunity, atherosclerosis and cardiovascular disease. BMC Med 2013;11:117.
- Herrington W, Lacey B, Sherliker P, et al. Epidemiology of atherosclerosis and the potential to reduce the global burden of atherothrombotic disease. *Circ Res.* 2016;118:535–546.
- Hansson GK, Robertson A-KL, Söderberg-Nauclér C. Inflammation and atherosclerosis. Annu Rev Pathol. 2006;1:297–329.
- Demopoulos CA, Karantonis HC, Antonopoulou S. Platelet activating factor—a molecular link between atherosclerosis theories. *Eur J Lipid Sci Technol.* 2003;105:705–716.
- Pearson TA, Mensah GA, Alexander RW, et al.; American Heart Association. Markers of inflammation and cardiovascular disease. *Circulation* 2003; 107:499–511.
- Palur Ramakrishnan AV, Varghese TP, Vanapalli S, et al. Platelet activating factor: a potential biomarker in acute coronary syndrome? *Cardiovasc Ther.* 2017;35:64–70.
- Cojocaru M, Cojocaru IM, Silosi I. Lipoprotein-associated phospholipase A2 as a predictive biomarker of sub-clinical inflammation in cardiovascular diseases. *Maedica (Buchar)* 2010;5:51–55.
- Triggiani M, Schleimer RP, Warner JA, et al. Differential synthesis of 1-acyl-2-acetyl-sn-glycero-3-phosphocholine and platelet-activating factor by human inflammatory cells. *J Immunol.* 1991;147:660–666.
- Prescott SM, Zimmerman GA, Stafforini DM, et al. Platelet-activating factor and related lipid mediators. Annu Rev Biochem. 2000;69:419–445.
- Marathe GK, Zimmerman GA, Prescott SM, et al. Activation of vascular cells by PAF-like lipids in oxidized LDL. *Vascul Pharmacol.* 2002; 38:193–200.
- Rainger GE, Chimen M, Harrison MJ, et al. The role of platelets in the recruitment of leukocytes during vascular disease. *Platelets* 2015; 26:507–520.
- Weyrich AS, McIntyre TM, McEver RP, Prescott SM, et al. Monocyte tethering by P-selectin regulates monocyte chemotactic protein-1 and tumor necrosis factoralpha secretion. Signal integration and NF-kappa B translocation. J Clin Invest. 1995;95:2297–2303.
- Prescott SM, McIntyre TM, Zimmerman GA, et al. Inflammation as an early component of atherosclerosis and vascular damage—a role for P-selectin and platelet-activating factor. Jpn Circ J. 1996;60:137–141.
- Lacasse C, Turcotte S, Gingras D, et al. Platelet-activating factor stimulates interleukin-6 production by human endothelial cells and synergizes with tumor necrosis factor for enhanced production of granulocyte-macrophage colony stimulating factor. *Inflammation* 1997;21:145–158.
- Handley DA, Arbeeny CM, Lee ML, Van Valen RG, et al. Effect of platelet activating factor on endothelial permeability to plasma macromolecules. *Immunopharmacology* 1984;8:137–142.
- Tsoupras A, Lordan R, Zabetakis I. Inflammation, not cholesterol, is a cause of chronic disease. *Nutrients* 2018;10:604.
- Antonopoulou S, Fragopoulou E, Karantonis HC, et al. Effect of traditional Greek Mediterranean meals on platelet aggregation in normal subjects and in patients with type 2 diabetes mellitus. J Med Food. 2006;9:356–362.
- Sukriti S, Tauseef M, Yazbeck P, et al. Mechanisms regulating endothelial permeability. *Pulm Circ.* 2014;4:535–551.
- Gaut JP, Heinecke JW. Mechanisms for oxidizing low-density lipoprotein. Insights from patterns of oxidation products in the artery wall and from mouse models of atherosclerosis. *Trends Cardiovasc Med.* 2001; 11:103–112.
- Lord an R, Nasopoulou C, Tsoupras A, et al. The anti-inflammatory properties of food polar lipids. In: Mérillon J-M, Ramawat KG, eds. *Bioactive Molecules in Food*. Switzerland: Springer International Publishing; 2018:1–34.
- Dentan C, Lesnik P, Chapman MJ, et al. Phagocytic activation induces formation of platelet-activating factor in human monocyte-derived macrophages and in macrophage-derived foam cells. Relevance to the inflammatory reaction in atherogenesis. *Eur J Biochem.* 1996;236:48–55.
- Antonopoulou SN, Karantonis HC, Fragopoulou E, et al. PAF, a potent lipid mediator. In: Ad T, ed. *Bioactive Phospholipids Role in Inflammation* and Atherosclerosis. Kerala, India: Tranworld Research Network; 2008:85–134.
- Zheng G-H, Xiong S-Q, Mei L-J, et al. Elevated plasma platelet activating factor, platelet activating factor acetylhydrolase levels and risk of coronary heart disease or blood stasis syndrome of coronary heart disease in Chinese: a case control study. *Inflammation* 2012;35:1419–1428.
- Cavallo-Perin P, Lupia E, Gruden G, et al. Increased blood levels of plateletactivating factor in insulin-dependent diabetic patients with microalbuminuria. *Nephrol Dial Transplant*. 2000;15:994–999.
- Kudolo GB, DeFronzo RA. Urinary platelet-activating factor excretion is elevated in non-insulin dependent diabetes mellitus. *Prostaglandins Other Lipid Mediat*. 1999;57:87–98.

- Chen H, Zheng P, Zhu H, et al. Platelet-activating factor levels of serum and gingival crevicular fluid in nonsmoking patients with periodontitis and/or coronary heart disease. *Clin Oral Invest*. 2010;14:629–636.
- Detopoulou P, Fragopoulou E, Nomikos T, et al. Baseline and 6-week follow-up levels of PAF and activity of its metabolic enzymes in patients with heart failure and healthy volunteers—a pilot study. *Angiology* 2013;64:522–528.
- Satoh K, Imaizumi T, Yoshida H, et al. Increased levels of blood plateletactivating factor (PAF) and PAF-like lipids in patients with ischemic stroke. *Acta Neurol Scand.* 1992;85:122–127.
- 29. Burke JE, Dennis EA. Phospholipase A2 structure/function, mechanism, and signaling. *J Lipid Res.* 2009;50 suppl:S237–S242.
- Nomikos T, Fragopoulou E, Antonopoulou S, et al. Mediterranean diet and platelet-activating factor; a systematic review. *Clin Biochem.* 2018;60:1–10.
- Huang F, Wang K, Shen J. Lipoprotein-associated phospholipase A2: the story continues. *Med Res Rev.* 2020;40:79–134.
- Tellis CC, Tselepis AD. Pathophysiological role and clinical significance of lipoprotein-associated phospholipase A<sub>2</sub> (Lp-PLA<sub>2</sub>) bound to LDL and HDL. *Curr Pharm Des.* 2014;20:6256–6269.
- Zalewski A, Macphee C. Role of lipoprotein-associated phospholipase A2 in atherosclerosis: biology, epidemiology, and possible therapeutic target. Arterioscler Thromb Vasc Biol. 2005;25:923–931.
- Silva IT, Mello APQ, Damasceno NRT. Antioxidant and inflammatory aspects of lipoprotein-associated phospholipase A2 (Lp-PLA2): a review. *Lipids Health Dis.* 2011;10:170.
- Shi Y, Zhang P, Zhang L, et al. Role of lipoprotein-associated phospholipase A2 in leukocyte activation and inflammatory responses. *Atherosclerosis* 2007;191:54–62.
- Steen DL, O'Donoghue ML. Lp-PLA2 inhibitors for the reduction of cardiovascular events. *Cardiol Ther.* 2013;2:125–134.
- Kohno M, Yokokawa K, Yasunari K, et al. Induction by lysophosphatidylcholine, a major phospholipid component of atherogenic lipoproteins, of human coronary artery smooth muscle cell migration. *Circulation* 1998;98:353–359.
- Garza CA, Montori VM, McConnell JP, et al. Association between lipoproteinassociated phospholipase A 2 and cardiovascular disease: a systematic review. *Mayo Clin Proc.* 2007;82:159–165.
- Hu G, Liu D, Tong H, et al. Lipoprotein-associated phospholipase A2 activity and mass as independent risk factor of stroke: a meta-analysis. *BioMed Res Int.* 2019;2019:1–11.
- Chung H, Kwon HM, Kim J-Y, et al. Lipoprotein-associated phospholipase A<sub>2</sub> is related to plaque stability and is a potential biomarker for acute coronary syndrome. *Yonsei Med J.* 2014;55:1507–1515.
- Perrot N, Thériault S, Rigade S, et al. Lipoprotein-associated phospholipase A2 activity, genetics and calcific aortic valve stenosis in humans. *Heart*. 2020;106:1407–1412.
- Lordan R, Tsoupras A, Zabetakis I. Phospholipids of animal and marine origin: structure, function, and anti-inflammatory properties. *Molecules* 2017;22:1964.
- Lordan R, Tsoupras A, Zabetakis I, et al. Forty years since the structural elucidation of platelet-activating factor (PAF): historical, current, and future research perspectives. *Molecules* 2019;24:4414.
- Zimmerman GA, Elstad MR, Lorant DE, et al. Platelet-activating factor (PAF): signalling and adhesion in cell-cell interactions. In: Nigam S, Kunkel G, Prescott SM, eds. Platelet-Activating Factor and Related Lipid Mediators 2: Roles in Health and Disease. Springer; 1996:297–304.
- 45. Detopoulou PD, Karantonis HC, Antonopoulou S. Mediterranean diet and its protective mechanisms against cardiovascular disease: an insight into platelet activating factor (PAF) and diet interplay. *Ann Nutr Disord Ther* 2015;2:1016.
- Xanthopoulou M, Kalathara K, Melachroinou S, et al. Wine consumption reduced postprandial platelet sensitivity against platelet activating factor in healthy men. *Eur J Nutr.* 2017;56:1485–1492.
- Nasopoulou C, Tsoupras AB, Karantonis HC, et al. Fish polar lipids retard atherosclerosis in rabbits by down-regulating PAF biosynthesis and up-regulating PAF catabolism. *Lipids Health Dis.* 2011;10:213.
- Panayiotou A, Samartzis D, Nomikos T, et al. Lipid fractions with aggregatory and antiaggregatory activity toward platelets in fresh and fried cod (*Gadus morhua*): correlation with platelet-activating factor and atherogenesis. J Agric Food Chem. 2000;48:6372–6379.
- Nasopoulou C, Gogaki V, Panagopoulou E, et al. Hen egg yolk lipid fractions with antiatherogenic properties. Anim Sci J. 2013;84:264–271.
- Koussissis SS, Hadzistavrou E, Kalyvas V, et al. PAF antagonists in food: isolation and identification of PAF antagonists in honey and wax. *Revue Francaise Des Corps Gras* 1994;5:127.
- Fragopoulou E, Detopoulou P, Nomikos T, et al. Mediterranean wild plants reduce postprandial platelet aggregation in patients with metabolic syndrome. *Metabolism*. 2012;61:325–334.
- Zia-ul-Haq M, Khan BA, Landa P, et al. Platelet aggregation and antiinflammatory effects of garden pea, Desi chickpea and Kabuli chickpea. Acta Pol Pharm. 2012;69:707–711.

- Poutzalis S, Anastasiadou A, Nasopoulou C, et al. Evaluation of the in vitro antiatherogenic activities of goat milk and goat dairy products. *Dairy Sci Technol.* 2016;96:317–327.
- Antonopoulou S, Semidalas CE, Koussissis S, et al. Platelet-activating factor (PAF) antagonists in foods: a study of lipids with PAF or anti-PAF-like activity in cow's milk and yogurt. J Agric Food Chem. 1996;44:3047–3051.
- Lordan R, Walsh AM, Crispie F, et al. The effect of ovine milk fermentation on the antithrombotic properties of polar lipids. J Funct Foods. 2019;54:289–300.
- 56. Lordan R, Walsh A, Crispie F, et al. Caprine milk fermentation enhances the antithrombotic properties of cheese polar lipids. *J Funct Foods*. 2019;61:103507.
- 57. Poutzalis S, Lordan R, Nasopoulou C, et al. Phospholipids of goat and sheep origin: structural and functional studies. *Small Rumin Res.* 2018;167:39–47.
- Prasad K. Reduction of serum cholesterol and hypercholesterolemic atherosclerosis in rabbits by secoisolariciresinol diglucoside isolated from flaxseed. *Circulation* 1999;99:1355–1362.
- Karantonis HC, Antonopoulou S, Demopoulos CA. Antithrombotic lipid minor constituents from vegetable oils. Comparison between olive oils and others. J Agric Food Chem. 2002;50:1150–1160.
- Koussissis S, Semidalas C, Antonopoulou S, et al. PAF antagonists in foods: isolation and identification of PAF in virgin olive oil. *Rev Fr Corp Gras* 1993;40:323.
- Tsoupras A, Fragopoulou E, latrou C, et al. In vitro protective effects of olive pomace polar lipids towards platelet activating factor metabolism in human renal cells. *Curr Top Nutraceutical Res* 2011;9:105–110.
- Choleva M, Boulougouri V, Panara A, et al. Evaluation of anti-platelet activity of grape pomace extracts. *Food Funct*. 2019;10:8069–8080.
- Papakonstantinou VD, Lagopati N, Tsilibary EC, et al. A review on platelet activating factor inhibitors: could a new class of potent metal-based antiinflammatory drugs induce anticancer properties? *Bioinorg Chem Appl.* 2017;2017:6947034.
- Saeed SA, Simjee RU, Shamim G, et al. Eugenol: a dual inhibitor of plateletactivating factor and arachidonic acid metabolism. *Phytomedicine* 1995;2:23–28.
- Weisenberger H, Grube H, Koenig E, et al. Isolation and identifiction of the platelet aggregation inhibitor present in the onion, *Allium cepa*. *FEBS Lett*. 1972;26:105–110.
- Apitz-Cstro R, Cabrera S, Cruz MR, et al. Effects of garlic extract and of three pure components isolated from it on human platelet aggregation, arachidonate metabolism, release reaction and platelet ultrastructure. *Thromb Res.* 1983;32:155–169.
- Tsuchiya H, Sato M, Watanabe I. Antiplatelet activity of soy sauce as functional seasoning. J Agric Food Chem. 1999;47:4167–4174.
- Sagesaka MY, Uemura T, Watanabe N, et al. A new glucuronide saponin from tea leaves (*Carnellia sinensis* var. *sinensis*). *Biosci Biotechnol Biochem*. 1994;58:2036–2040.
- Shah BH, Nawaz Z, Pertani SA, et al. Inhibitory effect of curcumin, a food spice from turmeric, on platelet-activating factor- and arachidonic acid-mediated platelet aggregation through inhibition of thromboxane formation and Ca2+ signaling. *Biochem Pharmacol.* 1999;58:1167–1172.
- Tzotzas T, Filippatos TD, Triantos A, et al. Effects of a low-calorie diet associated with weight loss on lipoprotein-associated phospholipase A2 (Lp-PLA2) activity in healthy obese women. *Nutr Metab Cardiovasc Dis.* 2008;18:477–482.
- Ntzouvani A, Giannopoulou E, Fragopoulou E, et al. Energy intake and plasma adiponectin as potential determinants of lipoprotein-associated phospholipase A(2) activity: a cross-sectional study. *Lipids* 2019;54:629–640.
- Hatoum IJ, Nelson JJ, Cook NR, et al. Dietary, lifestyle, and clinical predictors of lipoprotein-associated phospholipase A2 activity in individuals without coronary artery disease. Am J Clin Nutr. 2010;91:786–793.
- Nelson TL, Hokanson JE, Hickey MS. Omega-3 fatty acids and lipoprotein associated phospholipase A(2) in healthy older adult males and females. *Eur J Nutr.* 2011;50:185–193.
- Gajos G, Zalewski J, Mostowik M, et al. Polyunsaturated omega-3 fatty acids reduce lipoprotein-associated phospholipase A(2) in patients with stable angina. *Nutr Metab Cardiovasc Dis.* 2014;24:434–439.
- De Stefano A, Mannucci L, Tamburi F, et al. Lp-PLA2, a new biomarker of vascular disorders in metabolic diseases. Int J Immunopathol Pharmacol. 2019;33:2058738419827154.
- Zhuo S, Wolfert RL, Yuan C. Biochemical differences in the mass and activity tests of lipoprotein-associated phospholipase A2 explain the discordance in results between the two assay methods. *Clin Biochem.* 2017;50:1209–1215.
- 77. EndNote [Computer software]. EndNote X9. Clarivate; 2013.
- 78. Covidence [Computer software]. *Covidence Systematic Review Software*. Veritas Health Innovation.
- 79. Higgins JPT, Thomas J, Chandler J, et al. *Cochrane Handbook for Systematic Reviews of Interventions*. Version 6.2. West Sussex, England: Cochrane; 2021.
- Academy of Nutrition and Dietetics. Evidence Analysis Manual: Steps in the Academy Evidence Analysis Process. Chicago: Research, International and Strategic Business Development Team; 2016.

- Hernáez Á, Castañer O, Tresserra-Rimbau A, et al. Mediterranean diet and atherothrombosis biomarkers: a randomized controlled trial. *Mol Nutr Food Res.* 2020;64:2000350.
- Makariou SE, Elisaf M, Challa A, et al. No effect of vitamin D administration plus dietary intervention on emerging cardiovascular risk factors in patients with metabolic syndrome. J Nutr Intermediary Metab. 2019;16:100093.
- Shank ar N. A Randomized Comparative Study on the Effect of Fresh Coconut in a Balanced Yogic Diet on Anthropometric Biochemical Immunological and Psychological Parameters in Healthy Adults. Swami Vivekananda Yoga Anusandhana Sansthana; 2017. Available at: http://hdl.handle.net/10603/ 216121. Accessed August 20, 2021.
- Kim M, Song G, Kang M, et al. Replacing carbohydrate with protein and fat in prediabetes or type-2 diabetes: greater effect on metabolites in PBMC than plasma. Nutr Metab (Lond). 2016;13:3.
- Kim M, Jeung SR, Jeong TS, et al. Replacing with whole grains and legumes reduces Lp-PLA2 activities in plasma and PBMCs in patients with prediabetes or T2D. J Lipid Res. 2014;55:1762–1771.
- Wooten JS, Nambi P, Gillard BK, et al. Intensive lifestyle modification reduces Lp-PLA2 in dyslipidemic HIV/HAART patients. *Med Sci Sports Exerc.* 2013;45:1043–1050.
- Rizos CV, Liberopoulos EN, Tellis CC, et al. Combining rosuvastatin with sartans of different peroxisome proliferator-activated receptor-γ activating capacity is not associated with different changes in low-density lipoprotein subfractions and plasma lipoprotein-associated phospholipase A<sub>2</sub>. *Metab Syndr Relat Disord*. 2011;9:217–223.
- Karantonis HC, Fragopoulou E, Antonopoulou S, et al. Effect of fast-food Mediterranean-type diet on type 2 diabetics and healthy human subjects' platelet aggregation. *Diabetes Res Clin Pract*. 2006;72:33–41.
- Najjar RS, Moore CE, Montgomery BD. Consumption of a defined, plant-based diet reduces lipoprotein(a), inflammation, and other atherogenic lipoproteins and particles within 4 weeks. *Clin Cardiol.* 2018;41:1062–1068.
- Roberts CK, Ng C, Hama S, et al. Effect of a short-term diet and exercise intervention on inflammatory/anti-inflammatory properties of HDL in overweight/obese men with cardiovascular risk factors. J Appl Physiol (1985). 2006;101:1727–1732.
- Richard C, Couture P, Desroches S, et al. Effect of an isoenergetic traditional Mediterranean diet on the high-density lipoprotein proteome in men with the metabolic syndrome. J Nutrigenet Nutrigenomics. 2014;7:48–60.
- Seyedi S, Mottaghi A, Mirmiran P, et al. The relationship between dietary patterns and lipoprotein-associated phospholipase A2 levels in adults with cardiovascular risk factors: Tehran Lipid and Glucose Study. J Res Med Sci. 2020;25:3–3.
- Detopoulou P, Fragopoulou E, Nomikos T, et al. The relation of diet with PAF and its metabolic enzymes in healthy volunteers. *Eur J Nutr.* 2015;54:25–34.
- Hlebowicz J, Persson M, Gullberg B, et al. Food patterns, inflammation markers and incidence of cardiovascular disease: the Malmö Diet and Cancer study. J Intern Med. 2011;270:365–376.
- Chen CW, Lin CT, Lin YL, et al. Taiwanese female vegetarians have lower lipoprotein-associated phospholipase A2 compared with omnivores. *Yonsei Med* J. 2011;52:13–19.
- Lee KK, Fortmann SP, Varady A, et al. Racial variation in lipoprotein-associated phospholipase A<sub>2</sub> in older adults. *BMC Cardiovasc Disord*. 2011;11:38–38.
- Kowalska K, Socha E, Milnerowicz H. Review: the role of paraoxonase in cardiovascular diseases. Ann Clin Lab Sci. 2015;45:226–233.
- Schindhelm RK, van der Zwan LP, Teerlink T, et al. Myeloperoxidase: a useful biomarker for cardiovascular disease risk stratification? *Clin Chem.* 2009;55:1462–1470.
- Appay V, Rowland-Jones SL. RANTES: a versatile and controversial chemokine. *Trends Immunol.* 2001;22:83–87.
- Ivanova EA, Myasoedova VA, Melnichenko AA, et al. Small dense low-density lipoprotein as biomarker for atherosclerotic diseases. Oxid Med Cell Longev. 2017;2017:1273042.
- Rees K, Takeda A, Martin N, et al. Mediterranean-style diet for the primary and secondary prevention of cardiovascular disease. *Cochrane Database Syst Rev* 2019;3:CD009825.

- Schwingshackl L, Hoffmann G. Mediterranean dietary pattern, inflammation and endothelial function: a systematic review and meta-analysis of intervention trials. *Nutr Metab Cardiovasc Dis*. 2014;24:929–939.
- 103. Mayr HL, Thomas CJ, Tierney AC, et al. Randomization to 6-month Mediterranean diet compared with a low-fat diet leads to improvement in Dietary Inflammatory Index scores in patients with coronary heart disease: the AUSMED Heart Trial. *Nutr Res.* 2018;55:94–107.
- Kaur R, Kaur M, Singh J. Endothelial dysfunction and platelet hyperactivity in type 2 diabetes mellitus: molecular insights and therapeutic strategies. *Cardiovasc Diabetol.* 2018;17:121.
- Satija A, Hu FB. Plant-based diets and cardiovascular health. Trends Cardiovasc Med. 2018;28:437–441.
- Rombaut R, Camp JV, Dewettinck K. Phospho- and sphingolipid distribution during processing of milk, butter and whey. Int J Food Sci Tech. 2006;41:435–443.
- Lordan R, Tsoupras A, Mitra B, et al. Dairy fats and cardiovascular disease: do we really need to be concerned? *Foods* 2018;7:29.
- Lordan R, Zabetakis I. Invited review: the anti-inflammatory properties of dairy lipids. J Dairy Sci. 2017;100:4197–4212.
- Craddock JC, Neale EP, Peoples GE, Probst YC. Vegetarian-based dietary patterns and their relation with inflammatory and immune biomarkers: a systematic review and meta-analysis. Adv Nutr. 2019;10:433–451.
- US Department of Health and Human Services, US Department of Agriculture. Dietary Guidelines for Americans, 2020-2025. 9th ed. Washington, DC: DietaryGuidelines.gov; 2020.
- 111. Pub lic Health England. Government Dietary Recommendations. London, UK: Public Health England; 2016.
- 112. National Health and Medical Research Council. Australian Dietary Guidelines. National Health and Medical Research Council; 2013.
- Reidlinger DP, Darzi J, Hall WL, et al. How effective are current dietary guidelines for cardiovascular disease prevention in healthy middle-aged and older men and women? A randomized controlled trial. Am J Clin Nutr. 2015;101:922–930.
- 114. Castro-Acosta ML, Sanders TAB, Reidlinger DP, et al. Adherence to UK dietary guidelines is associated with higher dietary intake of total and specific polyphenols compared with a traditional UK diet: further analysis of data from the Cardiovascular risk REduction Study: supported by an Integrated Dietary Approach (CRESSIDA) randomised controlled trial. *Br J Nutr* 2019;121:402–415.
- 115. Tsimikas S, Willeit J, Knoflach M, et al. Lipoprotein-associated phospholipase A2 activity, ferritin levels, metabolic syndrome, and 10-year cardiovascular and non-cardiovascular mortality: results from the Bruneck study. *Eur Heart J.* 2009;30:107–115.
- 116. Hosseini B, Berthon BS, Saedisomeolia A, et al. Effects of fruit and vegetable consumption on inflammatory biomarkers and immune cell populations: a systematic literature review and meta-analysis. Am J Clin Nutr. 2018;108:136–155.
- 117. Hu FB, Rimm EB, Stampfer MJ, et al. Prospective study of major dietary patterns and risk of coronary heart disease in men. *Am J Clin Nutr.* 2000;72:912–921.
- 118. Fung TT, Willett WC, Stampfer MJ, et al. Dietary patterns and the risk of coronary heart disease in women. *Arch Intern Med.* 2001;161:1857–1862.
- Norde MM, Collese TS, Giovannucci E, et al. A posteriori dietary patterns and their association with systemic low-grade inflammation in adults: a systematic review and meta-analysis. *Nutr Rev.* 2021;79:331–350.
- Rizzo M, Berneis K. Low-density lipoprotein size and cardiovascular risk assessment. QJM. 2006;99:1–14.
- Gazi I, Lourida ES, Filippatos T, et al. Lipoprotein-associated phospholipase A2 activity is a marker of small, dense LDL particles in human plasma. *Clin Chem.* 2005;51:2264–2273.
- 122. Cowan SF, Leeming ER, Sindair A, et al. Effect of whole foods and dietary patterns on markers of subclinical inflammation in weight-stable overweight and obese adults: a systematic review. *Nutr Rev.* 2020;78:19–38.