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DOES COFFEE CONSUMPTION ALTER PLASMA LIPOPROTEIN(A) CONCENTRATIONS? A SYSTEMATIC REVIEW.

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ABSTRACT:

Coffee consumption alters plasma lipid and cholesterol concentrations, however, its effects on lipoprotein(a) (Lp(a)) have received little study. The aim of this PRISMA compliant systematic review was to examine the role of coffee on serum Lp(a).

This study was prospectively registered (PROSPERO 2015:CRD42015032335). PubMed, Scopus, Web of Science and Cochrane Central were searched from inception until 9th January 2016 to detect trials and epidemiological studies investigating the impact of coffee on serum Lp(a) concentrations in humans.

We identified six relevant publications describing nine experimental trials of various designs. There were a total of 640 participants across all studies and experimental groups. In short-term controlled studies, consumption of coffee, or coffee diterpenes was associated with either a reduction in serum Lp(a) of $\leq 11 \text{ mg/dl}$ (6 trials, 275 participants), or no effect (2 trials, 56 participants). Conversely, one cross-sectional study with 309 participants showed serum Lp(a) was elevated in chronic consumers of boiled coffee who had a median Lp(a) of 13.0 mg/dl (range 0-130) compared with consumers of filtered coffee who had median Lp(a) 7.9 mg/dl (range 0-144)

The effect of coffee on Lp(a) is complex and may follow a biphasic time-course. The type of coffee and the method of preparation appear to be important to determining the effect on Lp(a)

Keywords: cafestol, coffee, diterpenes, kahweol, lipoprotein(a).

No. of words: 209

INTRODUCTION

Coffee is a caffeine-containing beverage prepared as an aqueous extract of the beans of the *Coffea* plant. It is commonly consumed in Western society (Doepker et al. 2016). Previous meta-analyses have demonstrated associations between coffee consumption (particularly unfiltered coffee) and serum lipid concentrations (Jee et al. 2001). In particular, plasma concentrations of LDL-cholesterol and total cholesterol increase in a dose-dependent manner with exposure to coffee (Jee et al. 2001, Cai et al. 2012). Two diterpenes: kahweol and cafestol have been shown to be implicated in the lipid-modulating effects of coffee (Heckers et al. 1994, Weustenvanderwouw et al. 1994). These diterpenes are sometimes trapped by the paper filter used in some methods of coffee preparation. Scandinavian boiled coffee was shown to contain 3-4 mg of each diterpene per cup, compared with less than 0.1mg of each diterpene when the coffee was filtered (Urgert et al. 1995, Urgert et al. 1997). This helps to explain the observation that different methods for brewing coffee result in different effects on serum lipids (Dusseldorp et al. 1991).

Lipoprotein(a) (Lp(a)) particles consist of low-density-lipoprotein-like particles which are covalently bound to apolipoprotein(a) (Bos et al. 2014). Serum concentrations of lipoprotein(a) are positively correlated with cardiovascular risk (Kamstrup et al. 2009). Evidence from a study employing Mendelian randomization suggests that the link is causal (Kamstrup et al. 2009). A recent meta-analysis has demonstrated that elevated Lp(a) is an independent risk-factor for stroke (Nave et al. 2015). Low-fat diets that result in weight loss do not appear to result in alterations in plasma Lp(a) and two comprehensive reviews have concluded that the effects of diet on plasma Lp(a) concentrations are negligible (Puckey et al. 1999, Bos et al. 2014). Nevertheless, the well documented lipid-modulating effects of coffee, and the increasing recognition of Lp(a) as a risk factor for cardiovascular disease warrant investigation as to whether coffee can modulate plasma concentrations of Lp(a). It was our intention to carry out a systematic review and meta-analysis of studies of randomized controlled trials investigating the effect of coffee consumption on plasma Lp(a) concentrations in humans.

Our extensive and systematic literature search uncovered a limited, but interesting body of knowledge on this topic. There were insufficient randomised-controlled trials to perform a meta-analysis, so, instead we summarised in narrative format all the available evidence from studies in humans.

METHODS

Registration and search strategy

This PRISMA compliant study was prospectively registered (PROSPERO 2015: CRD42015032335). PubMed, Scopus, Web of Science and Cochrane Central were searched from inception until 9th January 2016. All fields were searched for the terms: (coffee OR "coffee" OR coffee* OR caffeine OR caffeine* OR "caffeine") AND (lipoprotein a OR LPa OR LP(a) OR lipoprotein(a) OR lipoprotein(a). Additionally, in the PubMed database, the terms were searched as MESH headers and all subheadings were included in the searches.

Inclusion and Exclusion Criteria

Inclusion Criteria

This systematic review included all studies in humans that examined the relationship between the consumption of coffee (or extracts of coffee) upon plasma concentrations of Lp(a). The PICOS strategy is outlined in **Table 1**. When results of a study were reported more than once, the most recent or complete article, or the one with the largest sample size, was included.

The following criteria were applied for inclusion:

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- Controlled trials or crossover trials which reported serum Lp(a) concentration at the baseline and completion and included coffee consumption (or abstinence) as an intervention (and studies from which these data were not reported but could be obtained from the study authors)
- Prospective cohort studies or other epidemiological studies which reported serum lipoprotein(a) concentrations and coffee consumption (and studies from which these data were not reported but could be obtained from the study authors)

Exclusion Criteria

Studies which were not conducted in humans were excluded. Studies which did not enable us to obtain sufficient information regarding Lp(a) were also excluded, except when that information could be obtained from study investigators.

Study Selection

All relevant articles were independently reviewed by two investigators (PP & MCS). The above inclusion and exclusion criteria were used to evaluate each article for selection into the systematic review. A third investigator (SU) was consulted to resolve study inclusion and exclusion discrepancies.

Data extraction

Eligible studies were reviewed and the following data were abstracted: first author's name; year of publication; country were the study was performed; study design; number of participants (divided into experimental groups where appropriate); details of coffee intervention; age, gender and body mass index (BMI) of the participants; baseline systolic and diastolic blood pressures; baseline TC, HDL-C, LDL-C and TG; baseline and (where appropriate) follow-up values of plasma concentrations of Lp(a). Studies reported their

results in a variety of units. Where the units for lipids given in the units mmol/l they were converted to mg/dL by multiplying by the following conversion factors (HDL-C, 38.61; LDL-C, 38.61; TC, 38.61; Triglycerides, 88.50). Data extraction was carried out by two investigators (PP & CS)

Quality Assessment

In order to assess the risk of bias in trials included in this review, the Cochrane Collaboration's tool for assessing risk of bias in randomized studies was used (Higgins et al. 2011, Higgins et al. 2011). Appropriate sections of this tool were completed for the one cross-sectional study. No trials, which met the inclusion criteria, were excluded from the systematic review on quality grounds.

RESULTS

Search results and trial flow

The flow of papers through the process is shown in **Figure 1**. Our searches found 945 papers. An initial screen of titles and abstracts was performed in order to remove articles, which were clearly irrelevant. After reading the full-texts of the remaining 121 papers, we identified 6 relevant papers (Urgert et al. 1996, Urgert et al. 1997, Strandhagen et al. 2003, Yukawa et al. 2004, Bukowska et al. 2006, Correa et al. 2013).

Description of studies

The characteristics of the studies and their participants and methods of the relevant papers we found are summarized in **Table 2.** The methods employed in the studies were extremely diverse The Quality assessment is shown in **Table 3**. The papers were published between 1996 and 2013 and included one relevant epidemiological study and five experimental papers describing nine trials of various designs. There were a total of 640

participants across all studies and experimental groups. Included in these figures are studies, which did not report the effect of coffee on Lp(a) quantitatively, but where that data was kindly provided by the authors. The studies included crossover and parallel group designs as well as trials in which participants were followed through a time course of coffee consumption and coffee abstinence. Interventions included boiled and filtered coffee and coffee diterpenes dissolved in oil. Comparators included abstinence from coffee, alternative methods of coffee consumption and placebo oil, or oil stripped of diterpenes. The effects of coffee consumption upon plasma Lp(a) are summarized in **Table 4**.

DISCUSSION

With respects to the methods employed, the studies were very heterogeneous. In studies where two blends of coffee were prepared, masking of participants to the blend was possible; in other circumstances masking the coffee intervention would have been extremely difficult and was not attempted. Nevertheless, it is unlikely that a participant's knowledge of their intervention would affect their plasma Lp(a) in a manner that would introduce bias. The difficulty of producing a placebo alternative to coffee may explain the paucity of randomized placebo controlled parallel group studies.

Urgert *et al.* published a paper that reported the results of four clinical trials(Urgert et al. 1997). They called these: Trial A, Trial B, Trial C and Trial D (Urgert et al. 1997). All were of relevance to this systematic review, and together provide information about the magnitude and direction of the effect of coffee on Lp(a), and also the components within coffee responsible for these effects. Trial B and Trial C were randomised placebo-controlled trials, Trials A and D had alternative study designs.

'Trial A' which was designed to compare the effects on Lp(a) of diterpene-rich unfiltered coffee with filtered coffee (Urgert et al. 1997). After a run-in period of four weeks in which

the participants drank filtered, coffee, they were randomised to receive 0.9 l/day (5 cups) of either filtered coffee or cafetiere coffee (Urgert et al. 1997). The concentrations of the diterpenes in the coffee were measured and translated in to daily doses (Urgert et al. 1997). Filtered coffee provided less than 1mg/day of each diterpene. Cafetiere coffee provided 38 mg/day cafestol and 33mg/day kahweol. Repeated measurements of Lp(a) were taken over time. Cafetiere coffee produced a fall in Lp(a) which was maximal at 8 weeks (1.5 mg/dL) and which stabilized at around 0.5 mg/dL between weeks 12 and 24 (Urgert et al. 1997). This time course may be of interest in explaining the results of an epidemiological study, described later, in which coffee consumption was associated with elevated Lp(a).

In Trial B, Urgert *et al.* performed a double-masked randomised-controlled trial in which 32 participants were randomised to receive 3g/day of either placebo oil (a 3:2 w/w mixture of sunflower oil and palm oil) or coffee oil which gave a daily dose of 85 mg of cafestol and 103 mg of kahweol(Urgert et al. 1997). The intervention was administered for four weeks, after which a statistically significant difference was found between the two groups, with respect to Lp(a) concentrations which were lower by a median of 5.3 mg/dL in the coffee oil group than in the placebo oil group (Urgert et al. 1997). Whilst these results seem to demonstrate a clear effect of coffee diterpenes on Lp(a), it should be noted that the daily doses of diterpenes are rather high, compared to that which might be expected from dietary coffee consumption. Another study reported in the same paper the authors found that 0.9 l of cafetiere coffee provided a dose of 38 mg cafestol and 33 mg kahweol (Urgert et al. 1997).

Also reported in the same paper was 'Trial C' which used very similar methods to 'Trial B' and was also conducted over four weeks (Urgert et al. 1997). The 36 participants were randomised to receive 2g/day of placebo oil, coffee oil (equivalent to a daily dose of 57 mg cafestol and 69 mg kahweol), or coffee oil that had been stripped of cafestol and kahweol (Urgert et al. 1997). Coffee oil reduced LP(a) concentrations by 3.1 mg/dL, an effect that was

not seen with placebo oil or stripped oil (Urgert et al. 1997). These trials, although small, provide evidence that diterpenes are responsible for the acute effects of coffee consumption upon Lp(a) (Urgert et al. 1997).

Further insight into the agent responsible for the acute Lp(a)-lowering effects of coffee was provided by Trial D (Urgert et al. 1997). Participants received either a mixture of cafestol (60 mg/day) and kahweol (48-54 mg/day) dissolved in placebo oil, or cafestol alone (61-64 mg cafestol/day and ≤ 1 mg/day kahweol). After a seven-week washout period, during which they took placebo oil, they were crossed-over to the other treatment group (Urgert et al. 1997). Cafestol alone produced a reduction in Lp(a) of 3.5 ±0.8 mg/dL (mean ±S.D.) compared with 3.9 ± 1.0 for the mixture. The changes from baseline were statistically significant, but the differences between the groups were not. This suggests that cafestol is the major diterpene involved in Lp(a) reduction observed with acute consumption of coffee (Urgert et al. 1997). The results of this trial are interesting, but should be treated with caution, because of the small number of participants (5 in each group), and because two participants in treatment groups were switched to placebo after having elevated alanine amino transferase which exceeded the safety limits defined by the investigators.

By combining data from all four of their randomised controlled trials. Urgert *et al.* made the interesting observation that the initial concentration of Lp(a) in an individual appears to influence the responsiveness of Lp(a) to coffee (or diterpene) treatment. After pooling the data, the investigators stratified participants into tertiles according to baseline Lp(a). Those with the highest initial values of Lp(a) saw the largest absolute reductions after treatment. Coffee or diterpenes treated participants in the highest baseline Lp(a) saw a median change in Lp(a) of -6.5 mg/dL, compared with control participants in the same Lp(a) tertile. For the middle Lp(a) tertile, the median difference was -3.3 mg/dL, and for the lowest tertile, -0.3 mg/dL (Urgert et al. 1997). Whilst Urgert *et al.* had found no effect of filter coffee upon Lp(a) (Urgert et al. 1997), a later study by Strandhagen and Thelle demonstrated an increase in Lp(a) after four weeks of consumption of 600 mml filter coffee per day (Strandhagen et al. 2003). The study consisted of two four-week periods of coffee consumption and two three-week periods of coffee abstention (Strandhagen et al. 2003). During both coffee consumption periods, Lp(a) values were reduced In the first period, the median difference was -11 mg/dl, in the second period it was -4 mg/dl (Strandhagen et al. 2003).The authors described the results as inconsistent, because there was no change in Lp(a) during the first abstention period, but a median increase of 15 mg/dl during the second abstention period. By comparison, total cholesterol increased during both the consumption periods and decreased during both the abstention periods (Strandhagen et al. 2003). Nevertheless, given the relatively small number of participants, the large variation in baseline Lp(a) levels between individuals, this would appear to be interesting evidence of a Lp(a)-lowering effect of filtered coffee (Strandhagen et al. 2003).

Also employing filtered coffee, Correa *et al.* conducted a randomised crossover trial designed to compare the effects of medium roast coffee and medium light roast coffee on lipids and other biomarkers (Correa et al. 2013). The twenty participants drank three or four cups daily of the first roast, before switching over to the other type. The diterpene concentrations of the coffee were measured and, concentrations of cafestol were substantially higher than those seen in other studies employing filtered coffee (Correa et al. 2013). Medium light roast provided 5.36 mg cafestol and 0.79 mg kahweol per 150 mg cup; medium roast provided 6.3 mg cafestol and 0.51mg kahweol per 150 mg cup. Mean coffee consumption was 462 ml/day, equivalent to a daily dose of cafestol of approximately 20mg (Correa et al. 2013). There were no statistically significant changes in plasma Lp(a) throughout the trial. However the small sample size of the trial may have rendered it underpowered to detect differences in Lp(a). It is also possible that the relatively low cafestol

dose in this trial may have been insufficient to have an effect on Lp(a), although the trial did show interesting differences in cholesterol and biomarkers of inflammation. Importantly this study demonstrates that diterpenes are not always retained by a paper filter (Correa et al. 2013).

In a randomised double-masked crossover trial, Bukowska *et al.* compared "natural unfiltered" coffee and coffee "modified by water and pressure extraction" with intervention periods of 28 days (Correa et al. 2013). The study included 36 healthy volunteers and compared Lp(a) before and after the intervention. The authors found no statistically significant differences in mean Lp(a) for either form of coffee (Lp(a) before 'modified form' coffee 32 ± 24 mg/dL, after 38 ± 26 mg/dL; before 'natural coffee 31 ± 27 mg/dL, after 32 ± 28 mg/dL). The study did, however show an increase in homocysteine in participants drinking the "natural unfiltered coffee", however the variance in baseline homocysteine was much smaller that for Lp(a), thus the trial may have been underpowered to detect changes in Lp(a) (Bukowska et al. 2006).

Yukawa conducted a study in 11 healthy male students in which participants drank 150 ml coffee three times per day for a week, preceded and followed by abstinence periods in which they drank only mineral water (Yukawa et al. 2004). The study aimed to investigate the effects of coffee on lipid metabolism and the oxidative modification of LDL-C. There were no differences between serum Lp(a) concentrations at the end of the baseline period $(25.1\pm 16.2 \text{ mg/dL})$, the end of the coffee consumption period $(23.2 \pm 11.4 \text{ mg/dL})$, and the washout period $(23.7 \pm 13.9 \text{ mg mg/dL})$ (Yukawa et al. 2004). It is likely that this trial was underpowered to detect differences in Lp(a) over the time period employed, however, statistically significant decreases in TC and LDL-C were observed (Yukawa et al. 2004). The authors suggested that the relatively high dose of coffee used in this study (150 ml three times

a day) may explain the fact that opposite effects of coffee on TC and LDL-C were seen here, compared to other studies (Yukawa et al. 2004).

Urgert *et al.* conducted a cross-sectional study comparing serum concentrations of Lp(a) in 150 habitual consumers of boiled coffee and 159 consumers of filter coffee (Urgert et al. 1996). Participants aged 40-42 years who reported drinking five or more cups of coffee per day were included in the analysis. Higher plasma concentrations of Lp(a) were found in consumers of boiled coffee (median 13.0 mg/dL; range 0-130 mg/dL) than in those who drank filter coffee (median 7.9 mg/dL; range 0-144 mg/dL). There was evidence of a doseresponse relationship between boiled coffee consumption and Lp(a). The subset of boiled coffee drinkers who reported consuming nine or more cups of coffee per day had a median Lp(a) concentration of 13.6 mg/dL compared with 11.7 mg/dL for those who drank fewer than nine cups. For filter coffee the values were 8.0 mg/dL; and mg/dL (Urgert et al. 1996). Despite the fact that these results seem to be in opposition to those reported in experimental studies, they are convincing because of the relatively large number of participants and because the results appear to show a dose-response relationship between coffee and Lp(a). These results cannot demonstrate causality, nor can they tell us whether the same chemical components of coffee are responsible for the short term reduction, and the long term elevation of Lp(a), however the fact that consumers of filtered coffee had lower Lp(a) than consumers of boiled coffee, suggests the responsible component may be trapped in a filter in the same way as the diterpenes have been in some studies.

In seeking to explain this result, the authors referred to previous observations that coffee increases serum alanine aminotransferase acutely. This marker is also elevated in liver disease (Weustenvanderwouw et al. 1994, Vanrooij et al. 1995). The investigators suggested, therefore, that in the short term diterpenes may disturb hepatocyte integrity, an effect which would be expected to result in reduced circulating Lp(a) (Gregory et al. 1994, Vanwersch

1994). Because normal serum concentrations of alanine aminotransferase were seen in this study, it was proposed that adaption occurs when coffee is consumed chronically. How and when this adaption occurs is unclear. This result is interesting in light of 'Trial A' described above, Urgert *et al.* reported maximal Lp(a) reduction after 8 weeks of consumption of boiled coffee, with a much smaller reduction from baseline seen thereafter (Urgert et al. 1997). The time-course demonstrated in that experiments supports the hypothesis that acute and chronic exposure to coffee may have different effects on Lp(a).

Clinical implications

No clinical recommendations can be made based upon the current evidence. The possible biphasic effect of coffee on Lp(a) mean that whilst coffee may have a short term beneficial effect in reducing Lp(a), in the longer term it may prove to be detrimental. Furthermore, seemingly beneficial effects of coffee in reducing plasma Lp(a) are likely to be counteracted by the effects of coffee consumption at increasing plasma total cholesterol and low-density-lipoprotein cholesterol which have been observed in most trials. Additionally, whilst elevated serum concentrations of Lp(a) are correlated with increased incidence of cardiovascular and cerebrovascular disease, the therapeutic benefit of Lp(a)-lowering is less well understood. Lp(a) should be more frequently measured and reported in clinical trials to enable us better to understand its prognostic importance, and to learn how it is affected by dietary and pharmacological interventions. Of the 106 papers selected for full-text screening but rejected for not reporting Lp(a), almost all reported numerous other lipid parameters.

Limitations

A limitation of this systematic review is the heterogeneity of study designs and interventions we included. Because of the small number of trials investigating the effects of coffee consumption on Lp(a), we included all types of study design which included humans. The number of participants in trials was generally very small. With respect to the intervention, coffee came from a variety of sources and multiple methods of preparation were employed. Therefore the results are hard to assimilate, and it was not possible to perform a meta-analysis. Despite the heterogeneity in reported methods of coffee preparations examined, there is a lack of data regarding decaffeinated coffee and coffee produced by automated coffee machines.

Heterogeneity of baseline serum concentrations of Lp(a) was noted within and between trials. The variability in this parameter is likely to increase the sample size required to demonstrate statistically significant changes with treatment. Additionally dietary interventions are harder to control than pharmaceutical intervention, adding another source of variability between participants. Thus trials which showed no effect of coffee on Lp(a) (Yukawa et al. 2004, Bukowska et al. 2006, Correa et al. 2013) or which showed an equivocal effect (Strandhagen et al. 2003) may have been underpowered with respect to Lp(a), despite being able to demonstrate changes in other parameters with baseline values which displayed less variance.

All the studies included in this systematic review relied on participants accurately reporting their dietary habits, or carefully following instructions regarding coffee preparation and consumption. This is a methodological weakness of any research investigating diet, however there is no reason to suppose that incorrect reporting by participants would systematically bias the study, rather than increasing variance in all groups.

The majority of the trials were not placebo controlled. Clearly it is clearly difficult to provide a placebo for coffee, without prior knowledge of the active Lp(a)-modifying agent. Even with this knowledge, it would be hard to produce a placebo whilst being certain that the

difference could not be detected by taste. Several of the trials could have been made more rigorous by parallel comparison of coffee-consuming groups and abstaining groups.

CONCLUSIONS

The effects of coffee consumption on plasma Lp(a) are complex and are likely to be affected by the baseline Lp(a) concentration, the source of the coffee, the method of preparation, the dose and the duration of consumption. There is a trend towards Lp(a)lowering effects of short-term consumption, with increased Lp(a) seen in chronic coffee drinkers. There is a need for more widespread reporting of Lp(a) in clinical trials.

ADDITIONAL INFORMATION:

This systematic review has been prepared within Lipid and Blood Pressure Meta-analysis Collaboration (LBPMC) Group (www.lbpmcgroup.umed.pl). The authors declare no competing financial interests.

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DECLARATION OF INTERESTS

The authors declare no competing financial interests.

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Table 1. Description of the PICOS	S criteria used to	o define the research	question
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Parameter	Description
Population	Humans, without any restrictions.
Intervention	Coffee consumption, ingestion of coffee-derived products, abstinence from coffee in habitual consumers
Comparator	Placebo or abstinence from coffee consumption.
Outcome	Change in plasma concentration of lipoprotein(a) after intervention
Study Design	All study designs in humans.

Study	Bukowska et	Correa et al.	Strandhagen et	Urgert et al.	Urgert et al.	Urgert et al.	Urgert et al.	Urgert et al.	Yukawa et
	al.		al.		Trial A	Trial B	Trial C	Trial D	al.
									(Yukawa et
									al. 2004)
Publication	2006	2013	2003	1996	1997	1997	1997	1997	2004
Year									
Location	Poland	Brazil	Sweden	Norway	The	The	The	The	Japan
					Netherlands	Netherlands	Netherlands	Netherlands	
Design	Randomised	Crossover	Controlled	Cross-	Randomised	Randomised	Randomised	Randomised	Controlled
	placebo	Clinical Trial	Study	Sectional	Controlled	Controlled	Controlled	Controlled	Study
	controlled			Study	Trial	Trial	Trial	Crossover	
	crossover trial							Trial	
Comparison	Natural coffee	Medium roast	Filtered coffee	Boiled coffee	Filtered coffee	Placebo oil vs	Placebo oil vs	Cafestol vs	Coffee vs
	vs pressure	coffee vs	vs abstinence	drinkers vs	vs Cafetiere	coffee oil	coffee oil v	cafestol &	abstinence
	extracted	medium light		filter coffee	coffee		'stripped oil.	kahweol	
	modified	roast		drinkers					
	coffee								

Table 2. Design of the studies selected for analysis and demographic characteristics and baseline parameters of participants.

Trial	4 weeks of first	1 week run-in; 4	2 x (3 weeks	NA	4 weeks filter	2 weeks	1 week placebo	2 x (2 week	1 week
Protocol	intervention;	weeks first	abstinence, 4		coffee; 24	placebo oil; 4	oil; 4 weeks	placebo oil; 4	baseline, 1
	28 day break; 4	intervention; 4	weeks		weeks	weeks	randomised	weeks	week
	weeks second	weeks second	consumption)		randomised	randomised	intervention; 4	randomised	coffee, 1
	intervention	intervention			intervention;	intervention;	weeks follow	intervention, 7	week
					12 weeks	4 weeks	up.	weeks follow	washout
					follow up	follow up		up)	
Inclusion	Healthy	Age 20 y to 65 y,	Inclusion	Recruited as	N/A	N/A	N/A	N/A	N/A
criteria	participants at	plasma	criteria were	part of the					
	age 28-55	cholesterol <240	age	Norwegian					
	years (50%	mg/dl∞, blood	range 30–65 y,	National					
	smokers). The	glucose <5.56	free of	Health					
	study was	mmol/L,	clinically	Screening in					
	conducted in	nonsmoker or	recognized	1992, a					
	the summer	former	chronic	population.					
	months to	smoker (>2 y),	diseases	Aged					
	avoid vitamin	alcohol	such as	40-42 years					
	deficiencies.	consumption less	cardiovascular	Subjects were					
		than one drink	diseases,	considered					

	per day, absence	cancer, renal	eligible if			
	of chronic	disorders, liver	they were			
	diseases, and no	disease and	healthy, did			
	use of regular	diabetes	not take			
	medication	mellitus. They	any			
		were not on	medication			
		antiepileptic	known to			
		or cholesterol-	affect liver			
		lowering drugs,	enzymes or			
		had been using	serum lipids,			
		coffee on a	and did not			
		regular basis	consume			
		for at least 5y	more than			
		and were	three			
		currently	alcohol-			
		nonsmokers	containing			
		(at least for the	beverages per			
		last 6 months)	day			
	1	1		1		

Source and	Two	Two	Not stated, but	N/A	Roodmerk	N/A	N/A	N/A	Arabica
type of	commercially	commercially	provided by		(Douwe				coffee
coffee	available	available blends	investigators to		Egberts) a				(Ajinomoto
	blends: natural	(80% Coffee	ensure		blend of				General
	coffee (MK	Arabica L. cv.	consistency		Arabica and				Foods,
	Cafe – 100%	Bourbon and			Robusta beans				Inc., Japan)
	Arabica)	20% C.							
	vs. modified	canephora cv.							
	coffee with	Robusta) of							
	60% less	caffeinated,							
	quantity of 2-	roasted, ground							
	methylisoborneol	coffee							
	(MK Cafe								
	Feelings; both:								
	MK Cafe,								
	Poland)								
Methods of	Natural coffee	Filtered	Filtered	N/A	Filtered or	Oil	Oil	Oil	Coffee
coffee	vs pressure				Cafetiere				dissolved
preparation	extracted								in boiling

	coffee								water
Dose of	3 x 180 ml	3-4 x 150 ml	600ml/day	NA.	Filtered coffee	Placebo oil	Placebo oil	Cafestol	150 ml
coffee	daily. Each	cups of coffee		Participants	(0.9 L/day)	(3g/day)	(2g/day)		three times
	serving	per day: mean		who					per day
	prepared with	$482 \pm 61 \text{ ml/day}$		habitually					Each
	13g ground			consumed			Coffee oil	Cafestol +	serving
	coffee			five or			(2g/day)	kahweol	prepared
				more cups of			Stripped oil		with 8g
				boiled coffee	Cafetiere	Coffee oil	(2g/day)		coffee.
				per day were	coffee (0.9	(3g/day)			
				compared	L/day)				
				with matched					
				filter coffee					
				consumers					
Daily	N/A	Approx 20	Not reported	N/A	<1 (filtered)	0 (placebo	0 (placebo)	61-64	N/A
Cafestol						oil)	57 (coffee oil)	(Cafestol)	
dose (mg)					38 (Cafetiere)	85 (coffee oil	Not reported	60 (cafestol	1
							(stripped oil)	plus kahweol)	

Daily	N/A	Approx 2.5	Not reported	N/A	<1 (filtered)	0 (placebo oil	0	0-1 (Cafestol)	N/A
Kahwol							69	-	
									-
dose (mg)					33 (cafetiere	103 (coffee	Not reported	48-56	
						oil)	(stripped oil)	(cafestol plus	
								kahweol)	
Participants	36	20	120 (first trial	150 (boiled)	24 (filtered)	16 (placebo	15 (placebo	10 (cafestol)	11
			period); 116			oil)	oil)		
			(second trial				15 (coffee oil)	-	
			period)	159 (filtered)	22 (cafetiere)	16 (coffee	16 (stripped	10 (cafestol	
						oil)	oil)	plus kahweol)	
Age (years)	42.7±5.8	49±9	48.6 (29-65)	41 ± 1	29 ± 10	22 ± 2	22 ± 2	24 ± 4	Range 21-
				(boiled)					31
				41 ± 1					
				(filtered)					
Male (%)	44	30	22	52.7 (boiled)	48.9	46.9	58.3	100	100
				55.3 (filtered)					
BMI	24.3±2.5	27.0±3.8	25.7 ± 3.4	25 ± 4	22 ± 3	22 ± 2	22 ±2	21 ±2	NS
((kg/m2)				(boiled)					

				25 ± 3					
				(filtered)					
SBP	N/A	110.2 ± 9.2	125.6 ± 17.3	N/A	N/A	N/A	N/A	N/A	N/A
(mmHg)				N/A					
DBP	N/A	70.5 ± 6.9	78.8 ± 11	N/A	N/A	N/A	N/A	N/A	N/A
(mmHg)				N/A					
TC (mg/dL)	226 ± 35	$186 \pm 23 \infty$	$201 \pm 36 \infty$	$231 \pm 42 \infty$	$189 \pm 27 \infty$	$174 \pm 19 \infty$	$174 \pm 28 \infty$	$186 \pm 35 \infty$	185 ± 18
	(Modified) ∞			(boiled)					
	221 ± 37			$219 \pm 41 \infty$					
	(Natural) ∞			(filtered)					
HDL-C	57 ± 11	$46 \pm 12 \infty$	$56 \pm 15 \infty$	N/A	$58 \pm 12 \infty$	$58 \pm 12 \infty$	$54 \pm 12 \infty$	58 ±15 ∞	57 ± 13
(mg/dL)	(Modified) ∞								
	53 ± 11			N/A					
	(Natural) ∞								
LDL-C	125 ± 34	$120 \pm 19 \infty$	N/A	N/A	$116 \pm 31 \infty$	$97 \pm 19 \infty$	$104 \pm 23 \infty$	$116 \pm 27 \infty$	122 ± 25
(mg/dL)	(Modified) ∞								

	127 ± 38 (Natural) ∞			N/A					
TG (mg/dL)	123 ± 62 (Modified) ∞	97 ± 35	$110 \pm 67 \infty$	$190 \pm 137 \infty$ (boiled)	97 ± 35∞	$89 \pm 27 \infty$	79 ± 27 ∞	71 ± 18 ∞	93 ± 31
	129 ± 64 (Natural) ∞			$170 \pm 110 \infty$ (filtered)					
Lp(a)	32 ± 24	22 ± 26	NS	NA	20.8 ± 22.3	25.9 ± 23.8	24.4 ± 23.4	13.9 ± 7.5	25.1 ± 16.2
(mg/dl)	(Modified)	(median = 11.5)			(median = 9.2)	(median =	(median =17.7)	(median	
					(filtered	17.2)	(placebo oil)	=11.5)	
					coffee)	(placebo oil)		(cafestol)	
	31±27				15.2 ± 19.9	29.1 ± 32.7	16.6 ± 16.6	13.9 ± 7.5	
	(Natural)				(median=9.8)	(median	(median =9.2)	11.5 (median)	
					(cafetiere	=14.9)	(coffee oil)	(cafestol +	
					coffee)	(coffee oil)	22.1 ± 25.5	kahweol)	
							(median =12.8)		
							(stripped oil)		

 $Values are expressed as mean \pm SD unless otherwise stated; \\ \infty values converted to units expressed here using http://www.endmemo.com/medical/unitconvert/$

Abbreviations: SD: standard deviation; SEM: standard error of the mean; BMI: body mass index; NA: not available; SBP: systolic blood pressure, DBP: diastolic blood pressure; TC: Total Cholesterol; LDL-C: lowdensity lipoprotein cholesterol; HDL-C: high-density lipoprotein cholesterol; TG, triglycerides. **Table 3:** Assessment of risk of bias in the included studies using a checklist based on the Cochrane Risk of Bias Assessment for Randomised

 Trials (with appropriate sections completed for the one cross-sectional study).

Author and	Sequence	Allocation	Blinding of	Blinding of	Incomplete	Selective outcome	Other potential
date	generation	concealment	participants and	outcome	outcome data	reporting	threats to
			personnel	assessment			validity
Bukowska <i>et al</i> .	TT	T	т	т	T	T	т
(2006)	0	0	L	L	L	0	L
Correa <i>et al</i> .	I	ŢŢ	т	Т	T	T	T
(2013)	U	U	L	L	L	U	L
Strandhagen et	NA	NA	I.	I.	I.	IJ	I.
al. (2003)				Ľ		C	L
Urgert <i>et al</i> .	NA	NA	NA	L	L	I	T
(1996)	1111		147 1	L	L	C C	L
Urgert <i>et al</i> .							
(1997)	U	U	L	L	L	U	L
Trial A							

Urgert <i>et al</i> .							
(1997)	U	U	L	L	L	U	L
Trial B							
Urgert <i>et al</i> .							
(1997)	U	U	L	L	L	U	L
Trial C							
Urgert <i>et al</i> .							
(1997)	U	U	L	L	Н	U	L
Trial D							
Yukawa <i>et al</i> .							
(2004)	NA	NA	L	L	L	U	L

L: low risk of bias; H: high risk of bias; NA: Not applicable; U: unclear risk of bias.

Study	Design	Intervention / Exposure	Lp(a) at baseline	Lp(a) at endpoint	Summary
			mg/dL	mg/dL	
			Mean ± S.D.	Mean ± S.D. unless	
			unless otherwise	otherwise stated	
			stated		
Bukowska et al. (Bukowska et	Randomised	Natural coffee	31 ± 27	32 ± 28	No effect of coffee on
al. 2006)	crossover trial	Pressure extracted coffee	32 ± 24	38 ± 26	Lp(a)
					No difference between
					groups
Correa et al. (Correa et al.	Crossover Clinical	Medium roast coffee	22 ± 26	22 ± 26	No effect of coffee on
2013)	Trial		11.5 (median)	14.0 (median)	Lp(a)
		Medium light roast coffee	22 ± 26	23 ± 29	No difference between
			11.5 (median)	13.9 (median)	groups
Strandhagen et al.	Controlled Study	Filtered coffee	NS	-11 (median, 1 st	Lp(a) reduction during first
(Strandhagen et al. 2003)				consumption)*	period of coffee

Table 4: Summary of the results of studies included in the systematic review

					-4 (median, 2^{nd}	consumption. Lp(a)
					consumption)	increase during second
			Abstinence from coffee	NS	$+2$ (median, 1^{st}	abstention period.
					abstention)	(*P<0.05)
					+15 (median, 2 nd	
					abstention)*	
Urgert et al. (Urgert et	al.	Cross-Sectional	Boiled coffee drinkers	N/A	13 (0-130)	Higher Lp(a) in boiled
1996)		Study			Median(range)	coffee drinkers ($P = 0.048$)
			Filter coffee drinkers	N/A	7.9 (0-144)	
					Median(range)	
Urgert et al. (Urgert	Trial A	Randomised	Filtered coffee		Change from	Lower Lp(a) in cafetiere
et al. 1997)		Controlled Trial		20.8 ± 22.3	baseline:	coffee drinkers than
				9.2 (Median)	$+0.2 \pm 0.8*$	filtered coffee drinkers
					+0.3 (median)	(*P<0.05)
			Cafetiere coffee		Change from	
				15.2 ± 19.9	baseline:	
				9.8 (median)	-2.0 ± 0.8	

					-0.9 (median)	
Urgert et al. (Urgert	Trial B	Randomised	Placebo oil		Change from	Lower mean and median
et al. 1997)		Controlled Trial		$25.9\ \pm 23.8$	baseline:	Lp(a) in consumers of
				17.2 (median)	$+1.1 \pm 0.9$	coffee oil than consumers
					+0.5	of placebo oil (**P<0.01)
			Coffee oil		Change from	
				29.1 ± 32.7	baseline:	
				14.9 (median)	-5.5 ± 1.4 **	
					-4.8 (median)**	
Urgert et al. (Urgert	Trial C	Randomised	Placebo oil		Change from	Lp(a) lowest in coffee oil
et al. 1997)		Controlled Trial		24.4 ± 23.4	baseline:	consuming
				17.7 (median)	-1.0 ± 1.6	group.(*P<0.05)
					+0.8 (median)	
			Coffee oil		Change from	
				16.6 ± 16.6	baseline:	
				9.2 (median)	-4.5 ± 1.3	
					-2.3 (median)*	

			Stripped oil		Change from	
				22.1 ± 25.5	baseline:	
				12.8 (median)	-1.1 ± 1.3	
					-0.3 (median)	
Urgert et al. (Urgert	Trial D	Randomised	Cafestol		Change from	Reduction in both groups
et al. 1997)		Controlled		13.9 ± 7.5	baseline:	compared to baseline
		Crossover Trial		11.5 (median)	$-3.5 \pm 0.8 **$	(**P<0.01)
					-3.1 (median)**	
			Cafestol & kahweol		Change from	
				13.9 ± 7.5	baseline:	
				11.5 (median)	$-3.9 \pm 1.0 **$	
					-3.5 (median)**	
Yukawa et al. (Yukawa et al.		Controlled Study	Coffee	25.1 ± 16.2	23.2 ± 11.4	No effect of coffee on
2004)						Lp(a)

FIGURE LEGENDS

Figure 1. Flow chart showing the number of studies identified, screened and included in the systematic review.