Effect of Coffee Consumption on Lipid Profile and Dyslipidemia Risk: Protocol for an Umbrella Review

Protocolo para una revisión de revisiones: impacto del consumo de café en el perfil lipídico y el riesgo de dislipidemia

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ABSTRACT

Introduction: Coffee is a drink that is associated with metabolic changes, including changes in the lipid profile. On the other hand, lipid alterations, such as increased LDL cholesterol or decreased HDL cholesterol, are associated with adverse cardiovascular outcomes. Taking into account the frequency of consumption of this drink and the recent evidence regarding its impact on cardiovascular and deaths, it is necessary to review the recent evidence to understand how coffee consumption modifies the lipid profile.

Methods and analysis: We searched Embase, Pubmed, BVS and Cochrane from inception to March 2021 with language restriction to French, Spanish and English. We included meta-analyses and systematic reviews that evaluated the impact of coffee consumption on the lipid profile in adults. Methodological quality of each study was evaluated using the Assessment of Multiple Systematic Reviews 2 (Amstar2) tool. The heterogeneity of the results reported using the I2 estimator was taken into account. A sensitivity analysis of the results was carried out by subgroups according to the quality of the included studies.

Keywords: coffee; caffeine; dyslipidemia; hypercholesterolemia; cholesterol.

RESUMEN

Introducción: El café es una bebida que se asocia con modificaciones metabólicas, entre ellas cambios en el perfil de los lípidos. Por su parte, los cambios lipídicos, como incremento de colesterol LDL o el colesterol HDL disminuido, se relaciona con desenlaces cardiovasculares adversos. Teniendo en cuenta la frecuencia de consumo de esta bebida y la evidencia sobre su impacto en el sistema cardiovascular y de muertes, es necesario comprender cómo el consumo de café modifica el perfil de los lípidos.
Métodos y análisis: Se llevó a cabo una búsqueda en Embase, Pubmed, BVS y Cochrane limitando por fechas desde la creación de las bases de datos, en francés, español e inglés. Se incluyeron metanálisis y revisiones sistemáticas que evaluarán el impacto del consumo de café en el perfil de lípidos en personas adultas. La calidad metodológica de cada estudio se evaluó mediante la herramienta “Assessment of Multiple Systematic Reviews 2” (Amstar2). Se tuvo en cuenta la heterogeneidad de los resultados reportados mediante el estimator $I^2$. Se llevó a cabo un análisis de sensibilidad de los resultados por subgrupos según la calidad de los estudios incluidos.

Palabras clave
café; cafeína; dislipidemia; hipercolesterolemia; colesterol.

Introduction

Hypercholesterolemia plays an important role in the development of cardiovascular disease (CVD), especially in the relationship between the amount of total cholesterol (TC), low-density lipoprotein (LDL) values, and the risk of cardiovascular events. People with hyperlipidemia have twice the risk of developing CVD compared to those with normal total cholesterol concentrations (1).

In turn, HDL cholesterol is a protective factor in CVD, while a low level of HDL, together with a high level of triglycerides (TG), may cause a higher incidence of CVD (2).

Coffee is a beverage with multiple effects on metabolism and it is estimated that around 3.5 billion cups of coffee are consumed worldwide every day (3). Several studies have found potential benefits with reduced risk of metabolic syndrome, obesity, and diabetes (4). Additionally, it has been found that moderate coffee consumption can have favorable effects in reducing cardiovascular mortality and CVD risk, among others (5,6).

There are several mechanisms by which coffee modifies the lipid profile. It has been described to have effects on lipogenesis, lipolysis, fatty acid $\beta$-oxidation, lipid transport, and fat digestion. These mechanisms are associated with the various components of the beverage, such as the neuromodulatory caffeine, which acts as an antagonist of the adenosine receptor as well as other components such as chlorogenic acids, trigonelline, or cafestol (4).

Among the effects of caffeine, it has been found to be associated with increased fat oxidation and glycogen mobilization in muscles, increased lipolysis and decreased body fat (7). Intake of green coffee bean extract, which provides 50 to 100 mg/day of chlorogenic acid, is associated with reductions in TC and LDL-C concentrations (between 8-10 and 3.5-5.5 mg/dL, respectively) in hypercholesterolemic subjects (8). For their part, the diterpenes cafestol and kahweol can increase the amount of total cholesterol in the blood by up to 30 mg/dL. This occurs because cafestol is a farnesoid X receptor agonist, which leads to an inhibition of bile acid synthesis and increased blood cholesterol levels. However, cafestol has also shown beneficial biological effects with antidiabetic, anticancer, and anti-inflammatory properties (4,9).

Some studies have found an unfavorable effect on the lipid profile, depending on the presentation and type of coffee consumed. In a randomized clinical experiment, it was found that coffee with caffeine had significant effects on the increase of LDL-C, TC and TG (7), although other publications affirm that the presence of caffeine does not modify the lipid profile variables (9).

On the other hand, compared to filtered coffee, unfiltered coffee has significantly increasing effects in patients with a history of hyperlipidemia on TC, LDL-C, and TG, and these effects increase with higher consumption (number of cups). This is considered to be a consequence of the presence of diterpenes, which are usually removed from the beverage when filters are used. Kahweol and cafestol increase the activity of cholesteryl ester transfer protein and phospholipid transfer protein while decreasing the activity of lecithin:cholesterol acyltransferase, thus contributing to an increase in LDL-C (7,9,10,11,12). In contrast, filtered coffee does not appear to modify serum lipids, but has been associated with an increased risk of metabolic syndrome (13).
Similarly, although espresso-type preparations have higher concentrations of diterpenes, the reduced portion size of the beverage decreases the amount consumed. Percolated and instant preparations have low levels of cafestol and kahweol, while French press preparations have the highest concentration of these diterpenes (9).

Considering that dyslipidemia is a cause of CVD (4) and that the literature on the impact of coffee on this condition is not conclusive regarding favorable and unfavorable findings, it is important to systematically review the best available evidence and identify the impact of the beverage on people's health.

Objectives

This review of systematic reviews of the literature aims to analyze, compare, and synthesize the evidence from available systematic reviews and meta-analyses on the effect of habitual coffee consumption on the development of dyslipidemia, and its relationship on the values of total cholesterol, LDL, HDL, and triglycerides, as well as the risk of developing dyslipidemia.

Methods

The review protocol was designed according to the guidelines of the Cochrane Collaboration and an adaptation of the Prisma-P checklist for systematic literature review and meta-analysis protocols (14).

Eligibility criteria consider systematic literature reviews and meta-analyses summarizing information regarding the effects of regular coffee consumption on the development of dyslipidemia and effects on total cholesterol, LDL, HDL, and triglyceride values. The inclusion and exclusion criteria are described in Table 2.

The primary outcomes will be: increase or decrease in TC values, LDL cholesterol, triglycerides, and HDL, and the development of dyslipidemias.

The literature will be searched in the Medline (via Pubmed), Embase, Cochrane Collaboration and LILACS (via BVS) databases. For this purpose, strategies will be designed that include controlled terms according to the base and other terms corresponding to the selected topics (Table 1) (15,16). Additionally, a secondary search will be carried out with a snowball strategy with the references cited in the manuscripts.

### Table 1.
Search strategy for the effects of habitual coffee consumption and dyslipidemia

<table>
<thead>
<tr>
<th>Subtopic</th>
<th>Terms related to coffee consumption</th>
<th>Terms related to outcomes</th>
<th>Terms related to type of publication and language</th>
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<tr>
<td>Dyslipidemia</td>
<td>Coffee OR “cafeté americano” OR “coffee consumption” OR “porción de café” OR “combinado de café” OR “café con leche” OR “caldo de café” OR “café de cama”</td>
<td>dyslipidemia OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia” OR “dyslipidemia”</td>
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<td>OR “coffee beverage” OR “coffee beverage” OR “coffee beverage” OR “coffee beverage” OR “coffee beverage” OR “coffee beverage” OR “coffee beverage” OR “coffee beverage”</td>
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</tbody>
</table>

Three reviewers will independently select manuscripts in two stages through Rayyan QCRI (17). In the first, the three independent reviewers will screen titles and abstracts; in the second, two independent reviewers will apply the criteria to pre-selected full-text manuscripts (Table 2). Discrepancies will be resolved by consensus. In addition, the references will be stored in the EndNote reference manager, and the duplication
of primary studies in the different systematic reviews will be taken into account and excluded.

Table 2. First stage of manuscript review

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Systematic literature reviews and meta-analyses that summarize information regarding the effect of regular coffee consumption on the development of dyslipidemia, and effects on lipid values; with no consumption or consumption at a lower dose of coffee. The studies must have been carried out in a population of men and women, adults over 18 years of age with or without a diagnosis of lipid profile alterations.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exclusion criteria</td>
<td>Manuscripts in English, Spanish, French, and Spanish will be considered. Review articles that have considered regular users of caffeine-contaminating drugs or beverages other than coffee that contain caffeine (tea, energy drinks, sodas, chocolate, etc.), or coffee derivatives such as green coffee extracts. Animal studies are also excluded.</td>
</tr>
</tbody>
</table>

Data collection and analysis

Data extraction will be carried out by three researchers following the format in Table 3. The quality of the selected documents will be evaluated using the Assessing Methodological Quality for Systematic Reviews 2 (AMSTAR 2) instrument (18).

Table 3. Format for data extraction

<table>
<thead>
<tr>
<th>Short reference</th>
<th>Author’s last name and initials of the first name(s). Year of publication and journal</th>
</tr>
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<tbody>
<tr>
<td>PICOC</td>
<td>Population, intervention, comparison and outcome</td>
</tr>
<tr>
<td>Searches</td>
<td>Databases, date and number of references. Date of last search</td>
</tr>
<tr>
<td>Gray Literature</td>
<td>Sources of unpublished literature consulted</td>
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<tr>
<td>Review quality</td>
<td>Quality scale used and level of quality reported.</td>
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<tr>
<td>Type and number of included studies</td>
<td>Primary studies of the review</td>
</tr>
<tr>
<td>Results and heterogeneity</td>
<td>Outcomes of interest in risk estimators (OR, RR, and HR) and their confidence intervals. Heterogeneity estimator (I^2 p for heterogeneity).</td>
</tr>
<tr>
<td>Potential biases of included publications</td>
<td>Publication, selection, participation, etc.</td>
</tr>
<tr>
<td>Conflict of interest</td>
<td>Reporting of conflicts of interest and funding sources</td>
</tr>
</tbody>
</table>

Ideally, information will be extracted from those publications with a score higher than 60% (more than 7 points); however, if no publications meeting the criteria are found, those with lower scores will be considered for inclusion, taking this into account when interpreting the results and proposing conclusions. Disagreements will be resolved by consensus and voting.

If discrepancies cannot be resolved by consensus, the kappa coefficient will be reported. The results of the systematic reviews will be summarized in tabular form for each of the outcomes of interest. The quality of the secondary studies will be presented in the tables and the heterogeneity of the results will be taken into account using the I2 estimator to assess the feasibility of calculating summary measures.

A sensitivity analysis of the results will be carried out by subgroups according to the quality of the included studies and by the dose of exposure to coffee consumption. Publication biases of the included secondary studies will be evaluated. Figures will be designed to facilitate the understanding of the relevant results.

Conflict of interest

The author’s team declares that they have no conflicts of interest.

References


