The Mediterranean Lecture: Wine and Thrombosis - From Epidemiology to Physiology and Back

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Abstract
The protective effect of moderate alcohol consumption on the risk of cardiovascular disease has been consistently shown in many epidemiological studies. Antiatherogenic alterations in plasma lipoproteins, particularly increase in high-density lipoprotein (HDL) cholesterol, are considered as the most plausible mechanism of the protective effect of alcohol consumption on coronary artery disease (CHD). Other potential mechanisms contributing to the cardio-protective effects of moderate alcohol consumption include anti-thrombotic down regulation of blood platelet function, as well as of the coagulation and fibrinolysis balance. Since the proposal of a "French paradox" in the early Nineties, the possibility that consuming alcohol in the form of wine might confer a protection against CHD above that expected from its alcohol content, has made the topic "wine and health" increasingly popular. Many epidemiological studies have explored such a possibility, by comparing specific alcoholic beverage types (wine, beer, liqueur) in respect to their relative capacity to reduce the risk of CHD. In parallel, experimental studies have been done, in which wine and wine-derived products have been tested for their capacity to interfere with molecular and cellular mechanisms relevant to the pathogenesis of CHD. Wine might indeed conceivably have other ethanol unrelated beneficial effects. The biological rationale for such a hypothesis has been linked to the enrichment in grape-derived, non-alcoholic components, that possibly make it peculiar in respect to other alcoholic beverages. In fact, while the mechanisms underlying the effects of alcohol on cardiovascular disease have been limited to lipid metabolism and the haemostatic system, those related to wine consumption have also been extended to specific anti-inflammatory, antioxidant and nitric oxide-related vaso-relaxant properties of its polyphenolic constituents. The effect of wine consumption has been carefully investigated to account for potential confounding of several conditions (inappropriate use of abstainers as control population, correlation between wine or total alcohol consumption and markers of healthy lifestyle and socioeconomic factors, diet, etc.). Strong evidence indicates that moderate wine consumption rather than confounders reduces both fatal and non fatal CHD events. In spite of the fact that the healthy effect of moderate intake of wine is by now well accepted, important issues remain to be resolved about the relationship between wine, alcohol and...
Introduction

Light to moderate drinkers of alcohol are at lower risk of vascular disease and death than non-drinkers [1-10]. The protective effect of moderate assumption of ethanol has been linked to anti-atherogenic alterations in plasma lipoproteins, as well as to anti-thrombotic modifications of blood platelet function, as well as of the coagulation and fibrinolysis balance [11]. The assumption of ethanol in the form of wine might conceivably have other non-ethanol related beneficial effects, linked to specific antioxidant and vaso-relaxant properties of its polyphenolic constituents [12].

We shall summarize here below experimental and epidemiological data supporting the cardio-protective effects of moderate wine intake.

Evidence from Experimental Studies

Anti-Atherogenic Effects

Alterations in plasma lipoproteins were considered as the most plausible mechanism of the protective effect of wine consumption on coronary artery disease (CHD) [13]. The antioxidant properties of polyphenols (grape-derived components of wine) may delay the onset of atherogenesis by reducing chemically mediated peroxidative reactions [14]. The potential in vivo antioxidant properties of wine have therefore been evaluated in terms of their ability to enhance plasma antioxidant status as well as to reduce specific oxidative processes. Studies in volunteers showed increased plasma antioxidant capacity or protection of LDL from oxidation after consumption of red wine [15-16]. A marked suppression of plasma post-prandial hydroperoxide increase was found in volunteers when a fatty meal was consumed with wine, instead of water [17]. Other relevant anti-atherogenic modifications concern the reported ability of red wine to increase plasma levels of HDL at levels above that obtained after alcohol consumption [12], or to reduce lipid deposition, in animal models, independently of their effects on lipid peroxidation [18]. As inflammation is a crucial component of the atherosclerotic process, it is of interest that resveratrol, a wine-derived polyphenol, significantly reduced the degree of colon injury, neutrophil infiltration, the levels cytokines and the COX-2 expression in an animal model of experimentally induced colitis [19].

Anti-thrombotic Effects

Other potential mechanisms contributing to the cardio-protective effect of moderate alcohol consumption include alterations of blood platelet function, coagulation and fibrinolysis. Alcohol seems to affect several factors involved in maintaining the delicate equilibrium between clot formation to protect against bleeding and clot dissolution to prevent blood clots from forming in arteries, and which have been implicated as risk factors for myocardial infarction [11].

Alcohol consumption has been associated with increased levels of tissue plasminogen activator (tPA), lower levels of fibrinogen and antithrombin III [20-22], and reduced platelet susceptibility to aggregation [23,24].

Polyphenols interfere with arachidonic acid metabolism, resulting in inhibition of platelet aggregation and reduced synthesis of pro-thrombotic and pro-inflammatory mediators [25-27]. Polyphenols can also down-regulate the expression of adhesive molecules and tissue factor activity, resulting in functional modulation of cell-cell interactions and procoagulant activities [26,28]. Tissue factor activity induced by any agonist was significantly reduced by resveratrol or quercetin [29-31]. Wine polyphenols were also shown to modulate nitric oxide (NO)-mediated responses. Fitzpatrick et al. [32] first showed that certain wine extracts were able to relax pre-contracted smooth muscle of intact aortic ring, and Andriambeloson et al. [33] demonstrated that the endothelium-dependent vaso-relaxation of rat aorta was mediated by an increase of NO aortic content. In addition, incubation of endothelial cells with red wines up regulated NO synthase mRNA and protein expression, and produced up to three times more bioactive NO than did control cells [34]. Martin et al. [35] showed that both delphinidin and anthocyanin (two polyphenols from red wine), inhibited endothelial cell apoptosis via NO pathway and regulation of calcium homeostasis. Wine itself may ameliorate endothelial dysfunction. In healthy volunteers acute smoking caused a reduction in flow-mediated dilatation (FMD), but simultaneous ingestion of either red wine or de-alco-
holized red wine with smoking prevented such a change in FMD, suggesting that wine can decrease smoke's harmful effect on endothelium [39]. Wine acutely improves endothelial function, measured as FMD, also in patients with coronary artery disease, although there was no detectable change in plasma polyphenol levels after wine consumption.

Platelet adhesion to damaged endothelium and subsequent platelet aggregation are major steps in both thrombosis and atherogenesis. Several studies have evaluated the possibility that alcohol consumption in the form of wine might have effect on human platelet function, additive to those already known to be exerted by alcohol. However, the antiplatelet activity of wine polyphenols observed in vitro [36, 37] and in experimental animal models [38], seems not to confer significant benefit, above that offered by alcohol itself, in human studies.

A European Collaboration on "Wine and Cardiovascular Disease"

The E.U.-supported FAIR Program included a collaborative project among several European partners aimed at evaluating the effects of red wine polyphenolic extracts (RWPE) on vascular tone, haemostatic system, oxidative processes, and plaque development [41]. Evidence was obtained that RWPE might improve vascular function mainly through NO-mediated mechanisms, interfere with haemostatic and oxidative processes involved in the progression of vascular damage, and modulate early events of atherosclerosis. A study in healthy Spanish volunteers showed a significant increase in HDL cholesterol levels and a decreased oxidation of LDL after moderate red wine ingestion (30 g daily for one month) as compared to the same amount of alcohol given as polyphenol-poor spirit [41]. A significant increase in HDL cholesterol levels and a decreased oxidation of LDL was observed both after red wine and gin ingestion (Table 1). A significant fall was seen, after red wine only, in the expression of several markers of cardiovascular risk, including the activity of MAC-1 and other adhesion molecules [42]. In the same group of volunteers it was found that TNF-alpha-induced adhesion of monocytes to endothelial cells was virtually abolished after wine consumption, but only partially reduced after gin [43]. Thus ethanol itself has potential beneficial effects on the cardiovascular system, mainly increasing HDL-cholesterol and decreasing oxidation of LDL. Wine may give additional benefits due to its greater antioxidant and anti-inflammatory effects.

Wine and the Mediterranean diet

Besides red wine, important sources of polyphenols are other beverages, foods and fruits. The source of different polyphenols may therefore vary in different populations [44]. On the other hand, the advantages of certain dietary habits with respect to the risk of CHD, have been extensively shown [45, 46]. High consumption of cereals, legumes, vegetables, and fruits, low consumption of red meat and dairy products, and the traditional use of olive oil, are the main characteristics of what is now considered as the healthy Mediterranean diet [47]. The observed correlation between wine and other foodstuffs such as fish and fruit and CHD mortality, suggest that the maintenance of traditionally Mediterranean lifestyle habits may contribute to decline in CHD mortality [47, 48]. Wine is the beverage of choice for Mediterranean populations and although it is not considered to be per se a component of the Mediterranean diet, its regular and moderate consumption may contribute to the lower rates of fatal and non fatal CHD observed in these populations [47].

Evidence from Epidemiological Studies

An inverse association between moderate alcohol consumption and CHD has been consistently shown in many epidemiological studies [1-10], but important issues remain to be clarified.

Differences among Different Alcoholic Beverages

Specific alcoholic beverages may have a different benefit. Renaud and de Lorgeril suggested wine intake as one possible explanation for the lower than expected CHD mor-
tality rates in France (the "French Paradox") [7], and many subsequent epidemiological studies have explored the French-paradox-generated hypothesis. We reviewed such studies in two meta-analyses based on 26 articles (including more than 200,000 persons) reporting comparisons between wine and beer and the risk of CHD [1]. The overall relative risk (RR) of wine drinkers in respect to non drinkers was 0.68 (95% confidence interval (95% CI): 0.59-0.77), whereas the protection associated with beer drinking was 10% lower (in absolute terms). In studies reporting dose-response curves, a clear non-linear dose-response curve was found between wine and vascular risk (Figure 1), but we failed to show any significant relationship between different amounts of beer intake and vascular risk. After the publication of our meta-analyses, three additional articles have been published. In the last report from the Framingham Study [8] about alcohol consumption and ischemic stroke, wine (but not beer or spirits) was related to a decreased risk of ischemic stroke (RR=0.8; 95%CI: 0.6 to 1.0). In a report [10] on mortality and alcohol consumption in a very large USA cohort (128 934 adults, 16 431 deaths in 20 years of follow up), protective effect of moderate alcohol intake against cardiovascular deaths was essentially restricted to women (20% reduced risk versus a non significant 10% reduction in men) that were more likely to drink wine. In contrast, the association of different types of alcoholic beverages with CHD risk evaluated in 38 077 USA males was strongest for beer and liquor, and weakest for wine [5].

The dose-response relationship between alcohol intake and rate of CHD has been traditionally depicted as a J-shaped curve [3], which means that non drinkers have higher risks than moderate drinkers, but similar or lower rates than heavy drinkers. In our meta-analysis [1] we confirmed the "J-shaped" curve since, after an initial decrease in the vascular risk by increasing amounts of wine, the curve reaches a plateau at higher intake, and tends to be reverted at the highest amounts explored (Figure 1). A maximum reduction was predicted at 750 ml/day, but statistical significance was only reached up to the amount of 150-200 ml/day, indicating that the amount of wine for which the balance between risk and benefit is the best in favor of wine is in the range 1 to 2 drinks a day, in agreement with international guidelines [49].

**Differences According to Clinical Phenotypes**

In subgroup analysis according to type of events, we found comparable protective effects of wine [1]. Wine intake was associated, indeed, with a decrease of both CHD (RR=0.71) and cerebrovascular disease (RR=0.43), and with both nonfatal (RR=0.71) and fatal vascular events (RR=0.49) (Table 1). Wine consumption was not significantly associated with myocardial infarction in the Health Professional Follow-Up Study [5] but it was so with ischemic stroke in Framingham Study [8]. The impressive study of Klatsky et al. [10] confirm the protective role of wine against total mortality. Quite recently, de Lorgeril et al [4] found a positive association of moderate wine intake with recurrences in survivors of a recent myocardial infarc-

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**Fig. 1.** Dose-response best fitting model for wine effect against combined vascular risk, obtained pooling "dose-response" curves in seven prospective studies (adapted from ref. 6, with permission)
tion, suggesting that moderate alcohol (wine) consumption may have positive effect in secondary prevention too.

**A Sex-related Effect of Wine?**

In our meta-analysis [1] we found that in studies enrolling only males, the protection offered by wine was small (13%) and not significant, in contrast with studies enrolling both genders (47% protection). In the Health Professional Follow-Up Study [5] conducted on USA males, the effect of wine was also small and not significant. An ad hoc analysis of the two USA studies included in our meta-analysis [1] where only males had been enrolled, showed no protective effect of wine. However in the five studies conducted in USA, but enrolling both sexes, overall relative risk reduction in favor of wine drinkers was 0.46 [2].

In the study of Klastky et al. [10] too the protection appeared to be confined to women. Thus, a possible sex difference in the protective effect of wine on vascular risk might exist and explain apparently controversial results in different epidemiological studies. However, in a meta-analysis of 12 studies comparing alcohol consumption (without any specific beverage distinction) in groups of males or females, we have lately failed to evidenciate any significant sex difference in the reduction of cardiovascular risk [50]. It is tempting to speculate that wine but not alcohol might exert a sex-related protection against cardio-vascular disease, but either experimental or clinical support to this hypothesis is still lacking. As a last comment in relation to the reported association between moderate alcohol consumption and increased risk of breast cancer [51], evidence suggests that the overall effect of moderate wine intake on women health may actually be favorable.

**Genetic Regulation of Alcohol Effects**

A strong interaction between alcohol intake and a common polymorphism in the gene for alcohol de-hydrogenase [52] has been observed; this finding supports causality of the polymorphism in the gene for alcohol de-hydrogenase [52].

**References**