Acute heavy alcohol intake increases silent myocardial ischaemia in patients with stable angina pectoris

Juhani Rossinen, Juhani Partanen, Pekka Koskinen, Lauri Toivonen, Markku Kupari, Markku S Nieminen

Abstract

Objective—To evaluate the effect of acute alcohol ingestion on myocardial ischaemia in patients with coronary heart disease and stable angina.

Design—Randomised crossover study using fruit juice with and without ethanol.

Setting—Division of cardiology in a university hospital.

Patients—20 patients with stable exertional angina and ≥ 50% luminal diameter narrowing of at least one major coronary artery.

Interventions—Each patient was studied on two separate days, once after administration of 1.25 g of ethanol per kilogram of body weight diluted to 15% in juice, and once after an equivalent volume of juice; both tests were in the evening and lasted 90 minutes. The patients were scheduled to have 8 periods of walking for 10 min according to a time table. An ambulatory electrocardiogram and the occurrence of anginal attacks were recorded and blood pressure and blood ethanol concentration were measured until the next morning.

Results—The blood ethanol concentration (mean (S.D.) range) rose to 28.8 mmol/l (13-3 (0-4)%). Alcohol raised the systolic blood pressure from 132 (16) to 141 (14) mm Hg (P < 0.05 compared with juice). The mean heart rate increased from 57 (7) to 64 (8) beats/min (P < 0.05) for 13 hours after ethanol ingestion compared with juice. The total duration of ischaemia during the ethanol test was 3.5 (median, range 0-80) min, compared with 0 (range 0-67) min for the juice test (P < 0.05). The difference resulted mainly from more silent ischaemia after ethanol ingestion (2-3 (0-80) v 0 (0-67) min; P < 0.05). The ST segment depression time integral increased during the ethanol test (4-4 (0-170) mm × min) relative to that during the juice test (0 (0-103) mm × min; P < 0.01) and especially during the following 13 hours after alcohol (3-5 (0-123) mm × min) compared with juice (0 (0-67) mm × min; P < 0.005). There were no changes in the number, duration, or ST segment depression time integral of the episodes of symptomatic angina, indicating that ethanol augmented the appearance of silent ischaemia.

Conclusion—Acute heavy alcohol drink-
ater narrowing of \( \geq 50\% \) in the left main coronary artery or luminal diameter narrowing of \( \geq 80\% \) in all three major epicardial vessels, ejection fraction \( \leq 25\% \) shown by left ventricular cineangiography, previous sustained ventricular tachycardia or fibrillation, the presence of other than sinus rhythm, or a history of alcohol abuse. We also excluded patients who used digitalis and those who had ST segment depression in the resting electrocardiogram or left ventricular hypertrophy, bundle branch block, or other conditions that might confound the evaluation of ischaemia.

The study was conducted in accordance with the Helsinki Declaration for Human Research and was approved by the ethics committee of the Department of Medicine of Helsinki University Central Hospital.

**METHODS**

After giving their informed consent, the patients were admitted to hospital and ambulatory electrocardiographic monitoring (Holter) was started at 1200 and finished on the next day at 1210. The patients were randomly assigned to having alcohol first or juice first in a crossover manner. The mean (SD) interval between the two tests was \( 7 (3) \) days (range 4–14). From 1700 to 1830 the patients were given 1·25 g of ethyl alcohol per kilogram body weight diluted to 15% in juice, yielding a total volume of 700 (126) ml, or an equivalent volume of the juice alone. The dose for a person whose weight is 70 kg is equivalent to 2·8 dl of 40% whisky or 9·8 dl of 11% wine calculated from volumes and was chosen to produce distinct intoxication. This made blind administration impossible. The patients were instructed to drink at an even pace over the 1½ hours. Their regular daily medications were continued throughout the study. Smoking was not allowed during electrocardiographic monitoring. The episodes of anginal pain and consumption of short-acting nitrates preparations were recorded. Blood samples were taken at 1700, 1900, and 0800 for alcohol concentration measurement by gas chromatography during the alcohol experiment and at the beginning of juice ingestion.

The patients had 10 minute exercise periods at 1200, 1400, 1600, 1800, 1900 and on the next day at 0800, 1000, and 1200 at which times they climbed 3 to 5 flight of stairs at rate that would normally provoke no angina. The length and duration of the exercises were kept constant for a patient during the whole study. The exercises during the ethanol test were accompanied by one investigator (JR).

The ambulatory electrocardiograms were recorded on tape with Marquette recorders (Marquette Electronics, Milwaukee, Wisconsin). V1 and V5-like leads were used.\(^{13}\) The electrocardiograms were analysed using the Marquette Series 8000 Holter analysis system without knowledge of the study phase or the patient’s identity. All recordings were analysed by the same investigator (JR). ST segment depression was measured at 80 ms after the J point in comparison with the level of the PR interval. In cases with a persistently depressed ST segment, this level was taken as baseline. An ST segment depression \( > 0·1 \) mV in amplitude compared with baseline and \( > 1 \) min in duration was regarded as an ischaemic episode. ST segment depression phases with an intervening period of \( > 1 \) min without ischaemia were calculated as separate episodes. The number of ischaemic episodes and the total duration of myocardial ischaemia and the ST depression time integral (magnitude times duration) were used to evaluate the ischaemic load. Ventricular premature complexes were detected by a semiautomatic template method (Marquette Electronics) and verified by the operator. The hourly averages of heart rate were obtained from the software report of the electrocardiographic scanner.

Blood pressure was measured using a sphygmomanometer when the Holter recording was started, at the beginning of drinking, at 1800, before and after the walking at 1900, and the next morning.

**STATISTICAL ANALYSIS**

The data are expressed as mean (SD) or median and range. Comparisons between the ethanol and juice tests were made with a Wilcoxon signed rank test or an analysis of variance with repeated measurements.\(^{14}\) A P value \( < 0·05 \) with the two-tailed test was considered statistically significant. A Kolmogorov-Smirnov one-sample test was used to assess whether the distribution of the data was normal. If not, a logarithmic transformation was made before the analysis of variance.

**Results**

The clinical characteristics of the patients are given in table 1 and their heart disease variables in table 2. Nineteen patients were taking \( \beta \) blockers and long-acting nitrates, 17 aspirin, 7 calcium-channel blockers, and 4 angiotensin converting enzyme inhibitors. Nine patients had arterial hypertension; none had diabetes.

The blood ethanol concentration was 28 (8) mmol/l (that is, 1·3 (0·4)%) two hours after

<table>
<thead>
<tr>
<th>Table 1 Clinical characteristics of the patients</th>
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<tr>
<td><strong>Characteristic</strong></td>
</tr>
<tr>
<td>Age (y)</td>
</tr>
<tr>
<td>Gender (F/M)</td>
</tr>
<tr>
<td>Weight (kg)</td>
</tr>
<tr>
<td>Duration of coronary heart disease (months)*</td>
</tr>
<tr>
<td>Number of previous myocardial infarctions</td>
</tr>
<tr>
<td>NYHA class:</td>
</tr>
<tr>
<td>II</td>
</tr>
<tr>
<td>III</td>
</tr>
<tr>
<td>Alcohol consumption (g of ethanol/week)*</td>
</tr>
<tr>
<td>(51 g equals 5·7 dl of wine)</td>
</tr>
</tbody>
</table>


<table>
<thead>
<tr>
<th>Table 2 Characteristics of the patients' heart disease*</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Characteristic</strong></td>
</tr>
<tr>
<td>Number of patients with 1 vessel/2 vessel/3 vessel disease</td>
</tr>
<tr>
<td>Average workload during the last 4 minutes of the exercise test (W)</td>
</tr>
<tr>
<td>Heart volume in chest x ray/body surface area (mL/m²)</td>
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<tr>
<td>Left ventricular ejection fraction (%)</td>
</tr>
<tr>
<td>Left ventricular end diastolic pressure (mm Hg)</td>
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</tbody>
</table>

*Values are numbers of patients or means (SD).
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The drinking period lasted from 1700 to 1830. The difference in heart rate between the ethanol and juice test was significant from 1800 to 0700 (P < 0.05).

Figure 1 Heart rate responses during the ethanol and juice tests (mean heart rates are shown). The drinking period lasted from 1700 to 1830. The difference in heart rate between the ethanol and juice test was significant from 1800 to 0700 (P < 0.05).

The difference in systolic blood pressure was significant from 1800 to 05:00 (P < 0.05).

Figure 2 Systolic blood pressure during ingestion of ethanol (black columns) and juice (hatched columns).

The mean heart rate between 1800 and 0700 was significantly greater during the alcohol test than during the juice test, 64 (8) compared with 57 (7) beats/min (fig 1). Heart rate was already increased with ethanol at 1800, an hour before taking blood for alcohol concentration measurement; and there was no indication of an effect on heart rate from the subsequent 0800 blood sampling during the alcohol test. The systolic blood pressure rose during ethanol ingestion (fig 2), but the diastolic blood pressure did not change significantly.

There were no differences between the juice and ethanol tests in blood pressure or heart rate before the drinking period.

The median number of ventricular premature complexes on the alcohol day was 4 (0–343) beats and 3.5 (0–632) beats on the juice day (NS).

Ethanol did not change the number of reported angina attacks (juice 0 (0–2) v

Table 3 Number of ischaemic attacks, episodes of angina, and consumption of short acting nitrates (median (range))

<table>
<thead>
<tr>
<th></th>
<th>Ethanol</th>
<th>Juice</th>
<th>Ethanol v juice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of painful episodes during Holter monitoring</td>
<td>0 (0–2) (mean 0.3)</td>
<td>0 (0–2) (mean 0.3)</td>
<td>NS</td>
</tr>
<tr>
<td>Number of ischaemic episodes during Holter monitoring</td>
<td>2 (0–17)</td>
<td>0 (0–24)</td>
<td>NS</td>
</tr>
<tr>
<td>Consumption of short acting nitrates during whole day</td>
<td>0 (0–2) (mean 0.3)</td>
<td>0 (0–2) (mean 0.3)</td>
<td>NS</td>
</tr>
<tr>
<td>Whole day:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total ischaemia (min)</td>
<td>3.5 (0–80) (mean 12.5)</td>
<td>0 (0–67) (mean 6.5)</td>
<td>P &lt; 0.05</td>
</tr>
<tr>
<td>Painful ischaemia (min)</td>
<td>0 (0–19) (mean 1.8)</td>
<td>0 (0–16) (mean 1.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Silent ischaemia (min)</td>
<td>2.3 (0–80) (mean 10.7)</td>
<td>0 (0–67) (mean 5.3)</td>
<td>P &lt; 0.05</td>
</tr>
<tr>
<td>1800–0700:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total ischaemia (min)</td>
<td>3.7 (0–57) (mean 10.3)</td>
<td>0 (0–42) (mean 4.5)</td>
<td>P &lt; 0.005</td>
</tr>
<tr>
<td>Painful ischaemia (min)</td>
<td>0 (0–19) (mean 1.6)</td>
<td>0 (0–16) (mean 1.0)</td>
<td>NS</td>
</tr>
<tr>
<td>Silent ischaemia (min)</td>
<td>2.1 (0–57) (mean 8.8)</td>
<td>0 (0–42) (mean 3.5)</td>
<td>P &lt; 0.01</td>
</tr>
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</table>
aggravation of ischaemia. Decreased pain perception, in accord with the analgesic effect of ethanol, may have helped to prevent patient awareness of the episodes.

Previous studies support our finding of increased silent ischaemia. Orlando et al.\textsuperscript{15} evaluated the effect of ethanol on exercise performance until the appearance of angina pectoris in patients with coronary artery disease in a study in which the achieved ethanol concentration was lower (1-0 (0-15)% vs) than in our study. Compared with control periods, the mean exercise time until angina was decreased after ethanol. Interestingly, there was more ischaemic ST segment depression after ethanol than after juice in the exercise level when angina was experienced, and the magnitude of ST segment depression increased with ethanol dose. This supports the hypothesis of increased silent myocardial ischaemia induced by ethanol. On the other hand, Pirwitz et al.\textsuperscript{16} have recently reported that intravenous ethanol induces epicardial coronary arterial vasodilation. In their study, however, the rate-pressure product did not increase, perhaps because the blood ethanol concentration achieved (0-86 (0-23)% vs) was lower than in our study. The increase in heart rate and blood pressure, and thus oxygen consumption, probably outweighs the possible vasodilator action of ethanol in the higher, rather toxic concentrations, used in our present study.

Ethanol depresses myocardial contractility in patients with heart disease.\textsuperscript{17,18} However, Greenberg et al. have observed a decrease in vascular resistance after ethanol that counterbalances the deterioration in cardiac performance.\textsuperscript{19} Our study gives no insight into whether left ventricular volume or myocardial contractility changed during ethanol ingestion. Thus it is also possible that by increasing left ventricular volume, ethanol might increase myocardial oxygen consumption.

In patients with ischaemic heart disease verified by coronary angiography, ST segment depression detected by ambulatory ECG monitoring is likely to indicate ischaemia.\textsuperscript{12} This suggests that the changes observed in our patients reflected a real myocardial oxygen deficit. The ST depression time integral is a valid indicator of ischaemia because the magnitude of ST segment depression in the Holter recording has been shown to correlate well with the severity of ischaemia demonstrated by an exercise test.\textsuperscript{20} Admittedly, the ST segment is affected not only by myocardial ischaemia, but also by ventricular hypertrophy, conduction disturbances, and pharmacological agents.\textsuperscript{11} Patients with these conditions were, however, excluded from the study.

All patients except one were taking β blockers. The mean rise of heart rate due to alcohol between 1800 and 0700 was only 7 (2) beats/min despite the high ethanol concentration (28 (8) mmol/l (1.3 (0.36)%)). The pulse frequency would probably have increased more after ethanol ingestion without β blockade,\textsuperscript{21} and thus ischaemia could have been even more marked without this treatment.

Moderate consumption of alcohol has been
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shown to reduce mortality in epidemiological studies. This contrasts with the harmful effects on ischaemia observed here. The daily ethanol dose associated with reduced mortality has, however, been much smaller than that used in the present experiment, which lead to signs of acute intoxication. Furthermore, there is only limited information on the influence of ethanol on mortality in patients with coronary heart disease. The study has potential limitations: day-to-day variation in myocardial ischaemia is considerable in patients with coronary heart disease. Our finding of ethanol-induced ischaemia, however, seems reliable because most of the increase in ischaemia occurred during ethanol intoxication and shortly after it. Although complete blinding is impossible in ethanol studies, the analysis of Holter recordings was blinded. The hospital ward does not represent the usual drinking circumstances of the patients and the mode of exercise performed can be questioned because quantification of the exercise level is difficult in floor climbing. It was, however, preferred over the conventional bicycle ergometry because it corresponds more closely to the exercise likely in usual drinking conditions. The amount of ethanol the patients consumed in this study was rather large and its relevance to lower doses is unknown: it should not be concluded from these results that social intake of 1 or 2 drinks causes ischaemia in patients with coronary heart disease. On the other hand, large amounts of alcohol are commonly drunk in Finland. Our results show that consuming alcohol in such amounts can provoke silent myocardial ischaemia in patients with coronary heart disease and should be discouraged.

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