Alcohol Reactions in Subjects of European Descent: Effects on Alcohol Use and on Physical and Psychomotor Responses to Alcohol

J. B. Whitfield and N. G. Martin

Self-reports of reactions to small amounts of alcohol, obtained between 1990 and 1992, were compared with reports of alcohol use, obtained in 1990–1992 and also in 1979–1981, in twin subjects of European descent. Data on subjective, physiological, psychomotor, and metabolic responses to a test dose of alcohol, taken in 1979–1981, were also available. Alcohol reactions were more common in women than in men, and were associated with less alcohol use, both at the time that information about reactions was obtained and as recorded on average 12 years previously, in both sexes. Physiological and psychomotor responses to alcohol were similar across the reaction groups, except that deterioration in standing steadiness was greater in those who subsequently reported adverse reactions to alcohol. Contrary to expectation, skin temperature changes after alcohol were less in the subjects who reported always reacting to alcohol than in the other groups. Subjective reports of intoxication were greatest in subjects who subsequently reported alcohol reactions. The pattern of twin pair concordance for reactions suggests low heritability, so alcohol reactions in subjects of European descent are not caused by a single gene of high penetrance of the type found in the Asian alcohol flush reaction.

Key Words: Alcohol Flush Reaction, Europeans, Alcohol Consumption, Skin Temperature, Alcohol Dependence.

The ALCOHOL flush reaction in Japanese, Chinese, and some other Asian subjects is a well-studied phenomenon that represents a unique example of a single gene having a major effect on alcohol use and alcohol dependence.1-3 This inherited aldehyde dehydrogenase-2 (ALDH2) deficiency is rare in non-Asian subjects; although similar reactions occur in some North and South American native groups they seem to be from a different enzyme defect.4 Reactions to alcohol in European-descent groups have attracted little attention and are, in the majority of cases, of unknown cause.

We have previously reported that acute reactions to small amounts of alcohol occur in ~5% of subjects of European descent, and that they have an aversive effect in that subjects reporting them also report lower alcohol consumption on a number of measures than nonreacting subjects. This may be a discrete metabolic phenomenon, like the Asian alcohol flush reaction caused by ALDH deficiency, but there is a need to define its effects more closely and to determine whether it shows the strong pattern of heritability to be expected if it is caused by an enzyme deficiency.

We have now completed a study of twin subjects previously tested with alcohol. This article describes the associations between self-reported alcohol reactions and alcohol consumption at different times in the subjects' lives, and symptoms of alcohol dependence. We have also tested whether physiological, subjective, and psychomotor responses to alcohol differ between European subjects who report reactions and those who do not. The concordance of twin pairs has been used to determine whether there is evidence for an inherited basis for European alcohol reactions.

Subjects and Methods

Three hundred and thirty-four subjects (157 males, 177 females) provided answers in 1990–1992 to a questionnaire containing sections on unpleasant reactions to small amounts of alcohol, alcohol use, and lifetime experience of symptoms or events associated with alcohol dependence. They were aged 27–46 years (mean = 34.1, SD = 4.8 years) at the time of participation and had also taken part, 9–13 years previously (1979–1981), in a study of alcohol metabolism and susceptibility to intoxication.4,5

All subjects were twins, but the number of complete pairs is rather less than half the number of subjects because it was not possible to contact or include both twins in every case. There were 29 pairs of monozygotic (MZ) and 25 pairs of dizygotic (DZ) male twins; 33 pairs of MZ and 33 pairs of DZ female twins; and 28 pairs of DZ twins of the opposite sex.

Questionnaire

All subjects completed a questionnaire with questions about reactions to small amounts of alcohol, quantity and frequency of alcohol use, number of drinks in the previous week, and lifetime occurrence of symptoms associated with alcohol use or dependence. The questions were:

Do you experience unpleasant reactions, such as flushing of the face or body, itching, drowsiness, or palpitations after drinking a small amount of alcohol (eg., one or two drinks):

Always
Sometimes
Never

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Please write in below the number that best describes how often (you) have had alcoholic drinks during the last 12 months.

Write the number that best describes how many drinks (you) usually have in a typical week.

On the chart below, please write the number of drinks you had on each day in the past week.

On average how many drinks would you have on each day that you have some alcohol?

Was there ever a time when:

- You drank too much
- You felt guilty about drinking
- Someone else objected to your drinking
- You were treated for a drinking problem
- You deliberately tried to cut down on your drinking, but were unable to do so
- You planned to stop drinking completely, but then failed to stick to your plan
- You got into physical fights while drinking
- You went on binges where you kept drinking for a couple of days or more without sobering up
- You went on drinking binges and neglected some of your usual responsibilities
- You got into trouble driving an automobile after drinking

Was there ever a time when your drinking had a harmful effect on:

- Your friendships and social life
- Your health
- Your marriage or home life
- Your work or employment opportunities

Have you ever had a drink first thing in the morning to steady your nerves or to get rid of a hangover?

Have you ever had drinking problems lasting at least a month?

A quantity-frequency score for alcohol use was calculated by multiplying the number of days of alcohol use per week and the number of drinks usually taken on a drinking day. A score for alcohol dependence was calculated by adding the number of positive answers to the dependence-related questions.

Data were also available for similar questions about alcohol consumption, but not alcohol reactions or alcohol dependence, which the subjects had answered some 12 years previously (in 1979–1981). Intake variables from that occasion were the usual number of drinks in a week and the quantity-frequency measure.

Alcohol Challenge Study

All subjects had taken part in an alcohol study, designed to investigate heritability of alcohol metabolism and intoxication, in 1979–1981. The amount of alcohol taken in that study was determined by the subjects' weight: 0.75 g of ethanol/kg. Baseline and postalcohol measurements of skin temperature, heart rate and blood pressure, performance on psychomotor tests, and subjective scores of intoxication had been recorded, and were available for analysis for effects of self-reported alcohol reaction status. Before drinking alcohol, the subjects had answered a questionnaire that included items on number of drinks taken per week, by type of drink, and their usual frequency and quantity of consumption. Skin temperature was measured by a thermistor probe taped to the cheek, and body sway was measured by the subject standing on a platform with a transducer that measured movement of the platform. Results reported in this article were obtained with the subjects' eyes open. Two questions were used to assess subjective perceptions of intoxication: they reported on their intoxication on a scale of 1–10 with 1 = completely sober and 10 = “The most drunk I have ever been.” And, they gave a yes/no answer to the question: “Would you drive a car now?” (yes was scored 1 and no was scored 0).

All of the subjects in the current study had been able to drink the alcohol in 1979–1981, over a period of 30 min. An unknown but small number of other volunteers had been unable to drink the alcohol, or had vomited, and no information from the initial study or follow-up was available for them.

Zygosity Determination

Details are given in Ref. 6. Twin pairs were classified as DZ if the twins were of the opposite sex, had differing types on blood grouping or plasma protein phenotyping, or differed markedly in height, natural hair type, or eye coloring. Otherwise, they were classified as MZ.

ADH Genotyping

Blood samples were available from 327 of the subjects who completed the follow-up questionnaire: 154 men and 173 women. Alcohol dehydrogenase (ADH)-2 and ADH3 genotypes were determined on DNA extracted from white blood cells. Appropriate sections of DNA, containing the areas that differ between ADH-1 and ADH-2, and between ADH-3 and ADH-2, were amplified by polymerase chain reaction using primers specific for ADH-2 or ADH-3. The primers used were: for ADH-2-5’ ATCTCTGAGATGTTGGcGT and 5’ GCCCTAAATTCAcAGGAAAG; for ADH-3-5’ CTTTAAAGATGAAGAAACT and 5’ CTCTTCCAGAGCCGACAGTC. After amplification, the ADH2 product was incubated with MaeIII that cuts the ADH2-2 sequence but not the ADH2-1, whereas the ADH3 product was incubated with SspI that cuts only ADH3-1. Digestion products were separated by electrophoresis and visualized with ethidium bromide to assign genotypes to the samples. Samples were also tested for ADH2-3 by allele-specific amplification, but no occurrences of ADH2-3 were found.

Statistical Methods

Subjects were classified into three groups (Always, Sometimes, or Never reacting) according to their answer to the question about alcohol reactions. Measured physiological, psychomotor, and subjective variables were tested for differences among reaction groups by analysis of variance. Alcohol use measures, and dependence scores, were transformed to log(x + 1) scores to reduce skewness; the constant value of 1 was added to allow subjects with 0 scores to be included. Tests for differences in the prevalence of reactions between men and women, and for association of alcohol reactions with ADH genotype, were performed by Fisher's exact test using StatXact Turbo 2.1 (Cytel Software Corporation, Cambridge, MA).

RESULTS

Prevalence

The proportion of men and women responding Always, Sometimes, or Never to the question about alcohol reactions is shown in Table 1. The prevalence of reactions differed significantly between men and women (p = 0.0001. Fisher's exact test). Because the prevalence of alcohol reactions, and the mean values for alcohol intake variables.

| Table 1. Reported Prevalence of Reactions in Males and Females |
|------------------|------------------|
|                  | Males            | Females         |
| Always           | 6 (3.6%)         | 12 (6.6%)       |
| Sometimes        | 30 (19.1%)       | 87 (37.9%)      |
| Never            | 121 (77.1%)      | 98 (55.4%)      |

Fisher's exact test, p = 0.0001.
Alcohol Use

Effects of alcohol reaction group on several measures of alcohol use are shown in Fig. 1. Alcohol intake measures were available at two times: at the time of the questionnaire (i.e., answers about reactions are contemporary with the answers about alcohol use) and 12 years previously (i.e., alcohol use recorded in 1979–1981 and reaction information provided later). The measures are of two types, based on number of drinks in a week or on quantity and frequency measures, and are shown separately for men and women. Alcohol reactions were associated with lower alcohol use, and their effect extended across the two occasions of questioning about alcohol use, some 10 years apart. Lifetime alcohol dependence scores did not differ significantly between the reaction groups, but there were few subjects with high scores (especially among the women) and few men who reported reactions.

Alcohol Metabolism

There was no significant association between blood alcohol concentrations, peak blood alcohol concentration, rate of decline, and the prevalence of reactions. The relationship between reactions and ADH2 or ADH3 type is not clear-cut; data are shown in Table 2. For ADH2 type, there was a significant association (p = 0.037, Fisher’s exact test), but the cell frequencies are low. Whereas the response Always is more likely in the ADH2-1.2 subjects, the response Sometimes is more common in the ADH2-1.1 group. ADH3 type was not significantly associated with alcohol reactions (p = 0.538).

Response to Experimental Alcohol Consumption

A number of physiological reactions to alcohol were measured in the 1979–1981 alcohol challenge study. For skin temperature, there were significant differences between alcohol reaction groups at times 1 and 2, but not at time 3 nor at (prealcohol) time 0. The Always group was significantly different from both the Sometimes group (p < 0.05), and the Never group (p < 0.01) at times 1 and 2. Skin temperature readings for the Sometimes and Never groups were essentially identical but, surprisingly, the Always group had the lowest temperature increase after alcohol.

For body sway, a marginally significant difference between groups (p = 0.059) was found at time 1. However, when the change in body sway between time 0 (prealcohol) and time 1 was calculated, there was a highly significant difference between the groups (F(2,33) = 7.29, p < 0.001). The increase in body sway in the Always group was significantly greater than for the Sometimes group (p < 0.05) or the Never group (p < 0.01); once again, the last two groups showed very similar results.

### Table 2. Subjects’ Responses to Alcohol Reaction Question, by ADH2 and ADH3 Types

<table>
<thead>
<tr>
<th>ADH2 Type</th>
<th>Always</th>
<th>Sometimes</th>
<th>Never</th>
</tr>
</thead>
<tbody>
<tr>
<td>ADH2-1.1</td>
<td>14 (4.7%)</td>
<td>90 (30.2%)</td>
<td>194 (65.1%)</td>
</tr>
<tr>
<td>ADH2-1.2</td>
<td>4 (13.8%)</td>
<td>4 (13.8%)</td>
<td>21 (72.4%)</td>
</tr>
<tr>
<td>p = 0.037, Fisher’s exact test</td>
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<td></td>
</tr>
<tr>
<td>ADH3-1.1</td>
<td>4 (4.4%)</td>
<td>24 (26.4%)</td>
<td>51 (57.7%)</td>
</tr>
<tr>
<td>ADH3-1.2</td>
<td>9 (4.9%)</td>
<td>52 (28.4%)</td>
<td>122 (66.7%)</td>
</tr>
<tr>
<td>ADH3-2.2</td>
<td>5 (9.6%)</td>
<td>17 (32.7%)</td>
<td>30 (67.7%)</td>
</tr>
<tr>
<td>p = 0.538, Fisher’s exact test</td>
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There were no significant differences (i.e., all \( p > 0.05 \)) for pulse rate, or systolic and diastolic blood pressures, between the alcohol reaction groups. Alcohol's effects on psychomotor test abilities, including reaction time, tracking ability, and divided attention tasks, did not differ significantly according to reaction group in either males or females, nor when results from both were taken together.

Effects of reported reactions on subjective impressions of intoxication, and on willingness to drive, showed several significant differences. The results for these two questions are shown in Figs. 2 and 3. There were significant effects of alcohol reactions on perceived intoxication at post-alcohol times 1 and 2, but not time 3, for both men and women. For men, the main difference was between those replying Always and those replying either Sometimes or Never, whereas for women the results for the Sometimes group were intermediate and significantly different from the Never group. For the willingness-to-drive question, reactions had little effect in women (who tended in any case to be more cautious than men in their response). In men, they had a significant effect at the later times (post-alcohol times 2 and 3) and once again only consistent (“Always”) reactions influenced the response. The Sometimes group showed negligible differences from the unaffected (“Never”) group.

Twin Concordance for Reported Alcohol Reactions

The pair concordances for reported alcohol reactions are calculated as polychoric correlations: MZ males = -0.02, MZ females = 0.39; DZ males = -0.08, DZ females = -0.34; and opposite-sex DZ pairs = -0.06. None of these are significantly different from 0. It was found that of 16 subjects who replied Always to the alcohol reaction question and whose cotwin’s response is also known, none had an Always reacting cotwin (regardless of zygosity) and more of their cotwins replied Never (11 subjects) than Sometimes (5 subjects).}

DISCUSSION

We would expect that people who find alcohol use unpleasant because of acute reactions to small amounts would
The existence of MZ twin pairs discordant for self-reported reactions shows that European alcohol reactions are not solely caused by a monogenic inborn error of metabolism, such as the well-explored Asian ALDH2 deficiency causing the alcohol flush reaction. ALDH deficiencies have been documented in a few subjects of European descent who experience alcohol reactions, but there are few well-studied examples and such enzyme deficiency seems to be an uncommon cause of alcohol reactions outside Asian groups.

A significant association between ADH2 type and the occurrence of reactions was found, but data suggest that this is a secondary factor, perhaps influencing the reaction enough to move some subjects from the “Sometimes” category to the “Always” category. Larger numbers of subjects would be needed to establish any ADH2 genotype effect in Europeans. However, it is of interest to note that ADH2 type, as well as ALDH2 type, affects the alcohol flush reaction in Asians. Thomasson et al. reported that among Chinese subjects with normal ALDH2 activity, ADH2-2 homozygotes had a greater increase in facial blood flow after alcohol than ADH2-1.2 heterozygotes. Data for ADH2-1 homozygotes was not reported; presumably there were insufficient subjects of this genotype in the group studied.

The physiological changes produced by alcohol are less than those reported in Asians, with no significant differences in pulse rate, blood pressures, or most tests of psychomotor performance between the reaction groups. Significant differences were found in facial skin temperatures, which would be expected to be the most prominent feature of flushing reactions, but these differences were in the opposite direction to what would be expected; those who reported “Always” reacting to alcohol had smaller skin temperature increases after alcohol. This also is in contrast with the results of Ward et al. who measured blood flow by a Doppler method and showed an increase in European flushers over nonflushers, although it was much smaller than the increase seen in an Oriental flusher.

Subjective assessments of intoxication after a test dose of alcohol were greater in the subjects who subsequently reported reactions. This was true both for the question about degree of intoxication (Fig. 2) and for the subjects' expressed willingness to drive a car (Fig. 3). There was also a significant difference in body sway between the subjects who reported reactions “Always” and those who did not, with a greater change after alcohol in those who reported alcohol reactions. Body sway, experienced as a feeling of unsteadiness, may well be one of the factors affecting subjects' perception of their degree of intoxication; it correlates significantly with self-report intoxication in both men and women in this sample even after controlling for the reaction responses (data not shown), and it is one of the aspects of the experience of intoxication claimed to influence subsequent alcohol dependence risk.

Perception and report of reactions are likely to be influenced by both physiological and psychological characteristics. People who drink little, for whatever reason, might have negative expectations about alcohol's effects and
therefore interpret their responses as unpleasant. However, average weekly alcohol consumption shows substantial heritability in both women and men and if the alcohol reactions (or the subjects' perceptions of them) are determined by alcohol consumption, then we would expect evidence of heritability for the reported reactions. This was not the case.

The way that these reactions are acquired, and the way that alcohol produces them, are still unknown. There may well be a number of causes for the self-reported reactions to alcohol in Europeans, including metabolic, physiological, and immunological (allergic) components. Some of the subjects may be found to have an inherited biochemical cause for their reactions, but these will be a minority among European subjects with alcohol reactions. Further exploration of the range of symptoms experienced in a much larger group of twin subjects, which is in progress, should shed light on heterogeneity of alcohol reactions.

REFERENCES