Familial Transmission of Substance Use Disorders

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**Background:** There is increasing evidence that substance use disorders are familial and that genetic factors explain a substantial degree of their familial aggregation. To perform a controlled family study of probands with several different predominant drugs of abuse, including opioids, cocaine, cannabis, and/or alcohol.

**Methods:** The subjects for the present study included 231 probands with dependence on opioids, cocaine, cannabis, and/or alcohol and 61 control probands, and their 1267 adult first-degree relatives. Diagnostic estimates were based on semistructured diagnostic interviews and/or structured family history interviews regarding each proband, spouse, and adult first-degree relative. The interview data were reviewed blindly and independently by clinicians with extensive experience in the evaluation and treatment of substance use disorders.

**Results:** There was an 8-fold increased risk of drug disorders among the relatives of probands with drug disorders across a wide range of specific substances, including opioids, cocaine, cannabis, and alcohol, which is largely independent from the familial aggregation of both alcoholism and antisocial personality disorder. There was also evidence of specificity of familial aggregation of the predominant drug of abuse.

**Conclusions:** Elevation in risk of this magnitude places a family history of drug disorder as one of the most potent risk factors for the development of drug disorders. These results suggest that there may be risk factors that are specific to particular classes of drugs as well as risk factors that underlie substance disorders in general.

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The familial aggregation of alcoholism has been well established. Although the results of numerous studies suggest that drug abuse is familial, there is a lack of adequate evidence from studies that use contemporary family study methods, including comparable control probands and direct interviews of relatives, in investigating the familial patterns of drug abuse.

Results: There was an 8-fold increased risk of drug disorders among the relatives of probands with drug disorders across a wide range of specific substances, including opioids, cocaine, cannabis, and alcohol, which is largely independent from the familial aggregation of both alcoholism and antisocial personality disorder. There was also evidence of specificity of familial aggregation of the predominant drug of abuse.

Conclusions: Elevation in risk of this magnitude places a family history of drug disorder as one of the most potent risk factors for the development of drug disorders. These results suggest that there may be risk factors that are specific to particular classes of drugs as well as risk factors that underlie substance disorders in general.

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The present study was designed to investigate the familial aggregation of drug and alcohol disorders across a broad range of drugs, including opioids, cocaine, mari-
SUBJECTS AND METHODS

SAMPLE CHARACTERISTICS

Probands

The sample of 299 probands was composed of a total of 149 probands with drug dependence (87 probands with opioid, 27 probands with cocaine, and 35 probands with cannabis dependence) and 89 probands with alcohol dependence, recruited from outpatient specialty clinics for substance disorders, and 61 control probands with no evidence of a lifetime history of any diagnosis in DSM-III-R. The controls were recruited through a random-digit dialing procedure in the greater New Haven, Conn, area. Study criteria were presented in detail by Merikangas et al.

Relatives

There were a total of 1267 first-degree relatives, primarily composed of parents and siblings (42% parents, 51% siblings, and 7% offspring). Overall, 57% of the relatives of probands who met study criteria were interviewed directly, either in person or by telephone (approximately 25% of the total number of relatives). Twenty percent of the first-degree relatives who refused direct interviews were assessed via informant reports. The interview rates did not significantly differ by proband diagnostic group and are in accordance with previous family studies of substance abuse.

OVERVIEW OF PROCEDURES

The procedures of the current study included the following: (1) interviewers with experience in substance abuse or clinical psychiatry interviewed relatives blindly with respect to the disorder of the index family member; (2) direct semistructured diagnostic interviews were applied, with expanded information on the correlates and course of patterns of drug use, particularly with respect to polysubstance use and abuse; (3) semistructured informant interviews were used, with expanded sections on substance abuse to obtain family history information; and (4) standardized diagnostic criteria based on information from both clinical material and informant reports were used in conjunction with the direct interview to assess diagnostic criteria for substance disorders and other psychiatric disorders (ie, best-estimate diagnoses).

Diagnostic Interview

The diagnostic interview for adults was the semistructured Schedule for Affective Disorders and Schizophrenia, current and lifetime versions, modified to obtain DSM-III and DSM-III-R criteria. With respect to substance abuse, the interview was modified to obtain more detailed information on the patterns of use of each drug class and their interrelationship, and on the course of alcohol and drug use and abuse.

Kappas derived from joint ratings of individual interviews were generally quite high for substance abuse (0.72-0.94) and somewhat lower for anxiety or affective disorders (0.54-0.78) across the first 3 series of training sessions. Comparison of diagnoses obtained through direct face-to-face interview and those from telephone interviews in both the present study and previous studies showed high levels of agreement across all diagnostic categories.

Family History Information

Family history information was obtained by means of a modified version of the Family History–Research Diagnostic Criteria developed by Andreasen et al for data collected by the family history method designed to obtain both DSM-III and DSM-III-R diagnoses in adults and children. The family history interview was modified to obtain more extensive information on patterns and sequelae of drug and alcohol use as well as extent of knowledge by the informant about the index subject.

Diagnostic Procedures

The probands were assigned to the 5 lifetime diagnostic substance disorder groups based on the predominant type of substance disorder through a blind and independent review by clinicians with extensive experience in the evaluation and treatment of substance abuse (K.R.M., S.S.O., B.J.R.). The predominant substance of abuse of probands and relatives was determined through a review of the following information on the life chart of substance use based on the diagnostic interview, treatment records where relevant, and family history information: age at onset, order of onset, quantity, frequency, chronicity, substance of choice, number of symptoms, and severity.

Statistical Analysis

To assess the demographic homogeneity of the proband groups and their relatives, χ² tests for contingency tables with discrete demographic variables and 1-way analyses of variance for continuous measures were applied. To account for familial clustering, mixed-effects multinomial logistic regression models were run with a program called MIXNO. MIXNO provides maximum marginal likelihood estimates of parameters for mixed-effects logistic regression of correlated nominal response data. In the present analyses, mixed-effects multinomial logistic regression models were used to investigate the association between (1) a 3-level substance disorder diagnosis variable (drug dependence ± alcohol dependence; alcohol dependence only; neither) in probands and relatives and (2) a 4-level predominant drug disorder variable (opioid, cocaine, cannabis, none) in probands (independent variable) and relatives (dependent variable), while adjusting for model covariates and correlated within family observations across all outcomes.

juana, and alcohol. Specifically, we examined the following patterns of familial associations: (1) the association between substance disorders in probands and relatives (ie, familial aggregation); (2) the association between drug disorders in probands and alcoholism in relatives and vice versa (ie, independence); and (3) the association between the predominant drug disorder in probands and relatives (ie, specificity).
Table 1. Demographic Distribution of Probands and Relatives

<table>
<thead>
<tr>
<th>Predominant Drug Disorder in Probands</th>
<th>Test for Homogeneity of Percentages or Means, P*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Opioid (n = 87)</td>
</tr>
<tr>
<td>--------------------------------------</td>
<td>----------------</td>
</tr>
<tr>
<td>Probands (n = 299)</td>
<td></td>
</tr>
<tr>
<td>Sex, % male</td>
<td>37</td>
</tr>
<tr>
<td>Mean ± SD age, y</td>
<td>35.3 ± 5.3*</td>
</tr>
<tr>
<td>Socioeconomic status,</td>
<td>82</td>
</tr>
<tr>
<td>Hollingshead Score &gt; I, %</td>
<td>39</td>
</tr>
<tr>
<td>Marital status, married or remarried, %</td>
<td>332</td>
</tr>
<tr>
<td>Relatives (n = 1267)</td>
<td></td>
</tr>
<tr>
<td>No.</td>
<td>332</td>
</tr>
<tr>
<td>Sex, % male</td>
<td>49</td>
</tr>
<tr>
<td>Mean ± SD age, y</td>
<td>49.2 ± 17.3</td>
</tr>
</tbody>
</table>

* Dunnett t test, df = 1; adjusted P < .05 for significant difference from controls.

RESULTS

DEMOGRAPHIC AND CLINICAL CHARACTERISTICS OF PROBANDS AND RELATIVES

Demographic characteristics of probands and relatives are presented in Table 1. Although the sex and age composition of the drug-abusing probands differed from that of the controls, there were no statistically significant differences in the age and sex distribution of the relatives.

There were no significant differences in the mean number of relatives of probands when compared with controls (ie, 4.5 relatives per proband). Of the 1267 first-degree biological relatives, 42% were parents, 51% were siblings, and 7% were offspring older than 18 years.

FAMILIAL AGGREGATION OF ALCOHOLISM AND DRUG DISORDERS

The familial aggregation of alcoholism and drug disorders is presented in Table 2 and Table 3. Table 2 presents the proportions of substance use disorders across broad categories of drugs, including “hard” drugs (ie, opioids, cocaine, stimulants, hallucinogens, or inhalants), “soft” drugs (ie, cannabis, sedatives, benzodiazepines, or barbiturates), alcohol, and nicotine among relatives by the predominant drug disorder of probands. Rates of hard drug disorders increased according to the deviance of the predominant drug among probands. Rates of abuse or dependence on hard drugs were greatest among the relatives of probands with opioid disorders (14.5%), moderately elevated among relatives with cocaine or cannabis disorders (9.6% or 8.4%, respectively), and lowest among relatives of probands with alcoholism (4.4%) when compared with relatives of controls (1.2%). In contrast, rates of alcoholism were increased only among the relatives of probands with cannabis or alcohol dependence compared with those of either the opioid or cocaine groups or the controls. Finally, rates of nicotine dependence were elevated among the relatives of all of the substance use disorder proband groups compared with those of controls. Approximately two thirds of the relatives reported a history of dependence on nicotine.

Table 3 presents the proportions of relatives with drug and/or alcohol disorders according to the presence or absence of drug and alcohol disorders in probands. The main effects of the familial aggregation of all alcoholism and drug disorders were highly significant: the relatives of probands with drug disorders had a 4.5-fold greater rate of drug disorders themselves than relatives of controls. Likewise, relatives of probands with alcohol disorders had a 2.0-fold greater rate of alcohol disorders than relatives of controls.

Statistically significant pairwise differences of proportions and their respective SEs, z scores, and P values are shown in the footnotes to Table 3. The proportions of relatives with both drug and alcohol disorders did not differ between those of probands with pure drug disorders (8.1%) and relatives of probands with comorbid drug disorders and alcohol disorders (9.1%). The lack of differences between rates of drug disorders among relatives of probands with comorbid drug disorders and alcoholism compared with relatives of probands with drug disorders only suggests that alcoholism and drug disorders are independent and that comorbidity is not an indicator of severity of substance disorders in probands.

Likewise, no significant evidence of coaggregation of pure forms of alcoholism and drug disorders emerged. Rates of pure drug disorders were increased among relatives of probands with pure drug disorders when compared with those of probands with alcohol only (P < .003); conversely, rates of pure alcoholism in the relatives were significantly elevated among relatives of probands with alcoholism only vs those with drug-only diagnoses (P < .001). Because similar patterns of familial aggregation of substance disorders emerged among relatives of probands in the drug and alcoholism group vs those in the drug-only group (see Table 3), these groups were collapsed in subsequent analyses.

Table 4 examines the association between drug and alcohol disorders in probands and relatives cross-classified according to the presence or absence of drug and alcohol disorders after controlling for demographic and clinical covariates. A mixed-effects generalized logit model, which allows for both adjustment for correlated observations (within families) and the simultaneous investigation of the 3 levels of the outcome variable (drug...
disorder with or without alcoholism; alcoholism alone; neither) was used. Applying the analyses presented in Table 4 to the subset of directly interviewed relatives yielded similar results. The intraclass correlations for familial clustering derived by the MIXNO program for nominal logistic regression analysis were 0.17 for alcoholism only vs none, and 0.04 for drug with or without alcoholism vs none.

After potential confounders were considered, these results were consistent with those in Table 3; alcohol and drug disorders aggregated independently in families, since there was no effect modification of the association between drug disorders in probands and relatives by the presence of alcoholism in the proband, nor was the association between alcoholism only in probands and relatives. The rates shown on the diagonal exceed the off-diagonal rates of all of the drug disorders and alcoholism. Inspection of the $z$ scores comparing rates of the predominant drug in probands and relatives yielded similar results. The intraclass correlations for familial clustering derived by the MIXNO program for nominal logistic regression analysis were 0.17 for alcoholism only vs none, and 0.04 for drug with or without alcoholism vs none.

The association between rates of the predominant drug disorder in probands and relatives is shown in Table 5. The statistically significant differences in the proportions between the predominant drug disorder in probands and the same predominant drug disorder in relatives are presented in the footnotes to Table 5, along with test statistics and $P$ values. The results in Table 5 show that there was a strong association between the predominant drug disorder in probands and relatives. The rates shown on the diagonal exceed the off-diagonal rates of all of the drug disorders and alcoholism. Inspection of the $P$ values of the $z$ scores comparing rates of the predominant drug in probands and relatives vs other proband drug groups showed significant levels of specificity for opioids, cannabis, and cocaine. Similar trends emerged when this analysis was applied to the subset of directly interviewed relatives.

Table 6 presents the results of the mixed-effects generalized logit model of the associations between the predominant drug disorders in probands and relatives after controlling for the effects of relevant covariates. The results in Table 6 confirm the findings of a direct association.
tion between the predominant drug disorders in probands and relatives reported in Table 5. For each of the specific drugs investigated, the adjusted odds ratio was greatest for the same drug disorder in the relatives (ie, 10.2 for opioids, 4.4 for cocaine, and 5.8 for cannabis) compared with the cross-drug associations. However, the overlapping and wide confidence intervals in the specific vs cross-drug comparisons also suggest that specificity should be considered as a trend rather than a conclusive finding.

The results of the present study show a remarkable degree of familial aggregation of substance abuse. There was an 8-fold increased risk of drug disorders among relatives of probands with drug disorders across a wide range of specific substances, including opioids, cocaine, cannabis, and sedatives, compared with that of relatives of controls. Elevation in risk of this magnitude demonstrates that a family history of drug abuse is one of the most potent risk factors for the development of drug abuse.

The increased risk of both drug disorders and alcoholism in male relatives across all proband groups is in accordance with the sex ratio for these disorders reported in large-scale epidemiological studies. The results of the present study confirm those of previous family studies that suggest independence of genetic factors predisposing to alcohol and drug disorders. In the present study there was no evidence of cross-transmission of drug disorders and alcoholism, nor did the presence of alcoholism modify the association between drug disorders in probands and relatives and vice versa.

The specificity of familial aggregation of the predominant drug disorder, particularly cannabis and alcohol, and to some extent the opioids, was unexpected. Several previous studies have shown a specific association between parental and offspring marijuana and nicotine use. Even more surprising in these earlier studies was the conclusion that parental and offspring resemblance was not attributable to direct exposure to parental substance use. These findings suggest that there may be some genes or environmental factors that predispose to the use of specific drugs rather than contributing to a deviant pattern of behavior or substance use in general.

In addition to some evidence for specificity of familial aggregation of the predominant drug disorder, the results also suggest that alcoholism and the specific drug disorders investigated herein appear to represent a continuum of severity. There was a direct increase in rates of serious drug disorders among relatives with increasing levels of “deviance” of substance disorders in probands ranging from alcoholism (4% drug disorders in relatives) to cannabis (8% in relatives) to cocaine (10% in relatives) to opioid abuse or dependence (15% in relatives) as compared with 1% among control relatives. Therefore, familial factors may be associated with increasing levels of progression along this continuum of severity of substance abuse rather than to the preference for a particular salient substance of abuse.

Familial clustering of drug abuse could be attributable to either common genetic or environmental factors that influence the development of drug disorders. Genetic factors could influence vulnerability to the development of drug abuse through individual differences in the effects of the drugs themselves, including metabolism, sensitivity, tolerance, side effects, and cognitive or psychological effects, or in alteration of affective, emotional, or cognitive states, such as reduction of stress, depression, or anxiety.

Families may convey an increased risk of substance disorders through both specific (eg, increasing environmental exposure to drugs or facilitating drug availability) and nonspecific (eg, impaired parenting behavior, exposure to marital discord, acute or chronic stress, negative life events, disrupted family structure, social deprivation, and physical, sexual, and emotional abuse) mechanisms. However, most of these family factors are not specific to drug disorders, since they also have been shown to characterize families of individuals with alcoholism as well as other psychiatric disorders.

There are several unique aspects of the design and methods of the present study that enhance its contribu-

Table 4. Adjusted Odds Ratios of Drugs and Alcohol Disorders in Relatives by Proband Diagnostic Group

<table>
<thead>
<tr>
<th>Relatives: Adjusted Odds Ratio (95% Confidence Interval)</th>
<th>Drug ± Alcohol</th>
<th>Alcohol Only</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Proband group</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Drug ± alcohol vs alcohol only</td>
<td>1.3 (0.9-2.2)</td>
<td>0.3 (0.2-0.6)</td>
</tr>
<tr>
<td>Drug ± alcohol vs controls</td>
<td>7.9 (3.6-17.4)</td>
<td>1.4 (0.8-2.5)</td>
</tr>
<tr>
<td>Alcohol only vs controls</td>
<td>6.0 (2.6-13.1)</td>
<td>4.2 (2.3-5.7)</td>
</tr>
<tr>
<td><strong>Covariates</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex of proband, female vs male</td>
<td>2.0 (1.3-3.2)</td>
<td>1.4 (0.9-2.1)</td>
</tr>
<tr>
<td>Antisocial in proband</td>
<td>0.8 (0.5-1.3)</td>
<td>1.4 (0.8-2.2)</td>
</tr>
<tr>
<td>Antisocial in relative</td>
<td>7.6 (4.1-14.0)</td>
<td>2.4 (0.5-1.4)</td>
</tr>
<tr>
<td>Sex of relative, female vs male</td>
<td>0.4 (0.2-0.6)</td>
<td>0.3 (0.2-0.5)</td>
</tr>
<tr>
<td>Age of relative, &lt;50 y vs &gt;50 y</td>
<td>8.2 (4.8-14.5)</td>
<td>0.8 (0.5-1.1)</td>
</tr>
<tr>
<td>Interview status</td>
<td>3.5 (2.2-5.7)</td>
<td>1.9 (1.3-3.0)</td>
</tr>
</tbody>
</table>

*P < .001.
†P < .01.
tion to current knowledge regarding familial factors and drug disorders. Design features include (1) a control group in which similar diagnostic and assessment methods were used; (2) probands with a range of drugs of abuse that strengthens the generalizability of the results; (3) a sufficient number of female probands to allow investigation of familial patterns of substance abuse among women; and (4) application of a statistical method that examines multinomial classification of probands and relatives simultaneously while controlling for the effects of clustered observations in families.

Limitations include (1) the lack of direct interview data on some of the relatives, necessitating the derivation of best-estimate diagnoses from family history information; (2) the imprecision of retrospective data in characterization of patterns of drug use and abuse (particularly among polysubstance abusers); (3) the lack of power to test adequately patterns of cross-transmission of specific drugs with such low base rates; and (4) possible selection biases with respect to clinical sampling of probands and recruitment of families with offspring, which may limit generalizability of the results.

Because the family has been shown to play a critical role in the etiology of drug use and abuse, it is important to obtain a deeper understanding of the complex mechanisms through which the family exerts its influence. Additional controlled family and twin studies are critical to identify patterns of expression of substance disorders and comorbid psychiatric disorders, to test the classic modes of genetic transmission of substance abuse, to determine the role of sex-specific patterns of transmission of substance abuse, and to elucidate the role of genetic and environmental factors and their interaction in its development.

Specificity of familial aggregation of some classes of drugs as well as independence from alcoholism suggests that research and treatment should focus on specific drugs of abuse and patterns of concurrent use to determine whether different genetic, biological, and environmental risk factors underlie their development.

<table>
<thead>
<tr>
<th>Predominant Drug Disorder in Relatives, %*</th>
<th>Opioid†</th>
<th>Cocaine‡</th>
<th>Cannabis§</th>
<th>Alcohol¶</th>
<th>All Drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Opioid</td>
<td>18.2</td>
<td>4.2</td>
<td>6.0</td>
<td>19.9</td>
<td>14.4</td>
</tr>
<tr>
<td>Cocaine</td>
<td>2.1</td>
<td>7.5</td>
<td>5.3</td>
<td>22.3</td>
<td>9.6</td>
</tr>
<tr>
<td>Cannabis</td>
<td>6.2</td>
<td>2.3</td>
<td>12.9</td>
<td>20.2</td>
<td>8.4</td>
</tr>
<tr>
<td>Alcohol</td>
<td>1.7</td>
<td>2.7</td>
<td>3.9</td>
<td>28.4</td>
<td>4.4</td>
</tr>
<tr>
<td>Controls</td>
<td>0.4</td>
<td>0.8</td>
<td>2.4</td>
<td>11.0</td>
<td>1.2</td>
</tr>
</tbody>
</table>

* For the proportion of relatives with the indicated specific drug disorder, statistically significant contrasts among proband groups, α = .05; P values are not adjusted for multiple comparisons. Boldface cells indicate specificity of concordance between probands and relatives.
† Opioid vs cocaine, percentage difference (% diff) = 8.1, SE = 2.2, z = 3.63, P < .001; opioid vs alcohol, % diff = 8.5, SE = 1.8, z = 4.78, P < .001; and opioid vs controls, % diff = 8.8, SE = 1.7, z = 5.86, P < .001.
‡ Cocaine vs controls, % diff = 6.7, SE = 2.8, z = 3.93, P < .001.
§ Cannabis vs opioid, % diff = 6.9, SE = 2.8, z = 2.44, P = .02; cannabis vs cocaine, % diff = 7.6, SE = 3.4, z = 2.22, P = .03; cannabis vs alcohol, % diff = 9.0, SE = 2.7, z = 3.34, P < .001; and cannabis vs controls, % diff = 10.5, SE = 2.7, z = 3.93, P < .001.
¶ Alcohol vs opioid, % diff = 8.5, SE = 3.1, z = 2.73, P = .006; alcohol vs cannabis, % diff = 8.2, SE = 3.7, z = 2.19, P = .03; and alcohol vs controls, % diff = 17.4, SE = 3.0, z = 5.86, P < .001.

| Predominant Drug Disorders in Relatives, Adjusted Odds Ratio (95% Confidence Interval)* |
|------------------------------------------|---------|---------|-----------|
| Predominant drug disorders in probands   | Opioids (n = 55) | Cocaine (n = 38) | Cannabis (n = 70) |
| Opioids (n = 87)                         | 10.2 (3.2-32.6)† | 2.8 (1.1-7.2)‡ | 2.4 (1.1-5.9)‡ |
| Cocaine (n = 27)                         | 2.2 (0.3-15.8)§ | 4.4 (1.5-13.2)§ | 1.8 (0.5-6.3) |
| Cannabis (n = 35)                        | 8.2 (1.7-39.1)§ | 1.9 (0.4-8.5) | 5.8 (2.6-12.9)† |
| Effects of confounders                   |          |         |           |
| Antisocial personality in proband       | 0.4 (0.2-1.2)  | 0.5 (0.2-1.7) | 1.0 (0.5-2.1) |
| Antisocial personality in relative      | 79.9 (26.9-237.4)‡ | 7.0 (1.4-35.4)‡ | 7.4 (2.3-23.4)‡ |
| Sex of proband, female vs male          | 1.1 (0.4-3.0) | 2.8 (1.1-7.0)‡ | 1.4 (0.7-2.6) |
| Sex of relative, female vs male         | 0.7 (0.3-1.7)‡ | 0.4 (0.1-0.9)‡ | 0.5 (0.3-0.9)‡ |
| Age of relative, <50 y                   | 8.1 (3.1-21.1)† | 49.7 (2.9-86.1)§ | 12.1 (4.5-32.7)† |
| Interview status of relative             | 0.4 (0.1-1.7)† | 0.9 (0.3-3.1) | 2.5 (1.3-4.7)§ |

* Boldface cells indicate specificity of concordance between probands and relatives.
† P < .001.
‡ P < .05.
§ P < .01.
In terms of prevention, these findings support a shift in drug prevention programs from the universal level to programs that target the families of substance abusers.

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REFERENCES