Modeling Genetic and Environmental Influences in the Etiology of Conduct Disorder: A Study of 2,682 Adult Twin Pairs

Wendy S. Slutske, Andrew C. Heath, Stephen H. Dinwiddie, Pamela A. F. Madden, and Kathleen K. Bucholz
Washington University School of Medicine

Michael P. Dunne, Dixie J. Statham, and Nicholas G. Martin
Queensland Institute of Medical Research

The etiology of conduct disorder (CD) was examined retrospectively in a sample of 2,682 male, female, and unlike-sex adult twin pairs from the community-based Australian Twin Register. Model-fitting analyses indicated a substantial genetic influence on risk for CD, accounting for 71% of the variance (95% confidence interval [CI] = 32–79%). There was not a statistically significant effect of the shared environment in the best-fitting model of CD, but a modest effect of the shared environment on the risk for CD could not be rejected (95% CI = 0–32%). The magnitude of genetic and environmental influences for CD liability did not vary significantly for boys and girls, and specific genetic and environmental mechanisms important for the development of CD appeared to be largely the same for both sexes. The fit of a multiple-threshold model raises the possibility that CD may not necessarily be a discrete entity but rather an extreme of the normal variation in conduct-disordered behavior found in the general population.

Conduct disorder (CD) is one of the more prevalent childhood psychiatric disorders and one of the most common reasons that children are referred for psychiatric treatment (Kazdin, 1987; Robins, 1991). Children with CD, in addition to being more likely to suffer from other childhood psychiatric disorders (Anderson, Williams, McGee, & Silva, 1987; Cohen et al., 1993), are much more likely than other children to suffer from alcohol and drug dependence, depression, anxiety disorders, and antisocial personality disorder (ASPD) as adults (Kazdin, 1987; Robins, 1978b; Robins & Price, 1991; Rutter, 1995). In fact, childhood CD may be one of the strongest predictors of psychopathology in adulthood.

Research and theory on the etiology of CD have emphasized the importance of the role of the family. In particular, a constellation of interrelated parental attributes such as ineffective parenting (Patterson, De-Baryshe, & Ramsey, 1989), poor parental supervision and discipline (Frick et al., 1992; Loebel & Dishion, 1983), parental conflict and divorce (Amato & Keith, 1991; Emery, 1982), and parental separation (Fergusson, Horwood, & Lynskey, 1994; Pearlson, Jalongo, Hunter, & Kellam, 1994) have all been associated with childhood antisocial behaviors. These risk factors may reflect aspects of the child's social environment or, alternatively, because of their correlation with parental psychopathology, may reflect the child's genetic risk for antisocial behavior (Robins, 1978a; also see Lykken, 1995). Lahay, Hardtgen, et al. (1988) found evidence to suggest that the association between parental divorce and offspring CD is due to parental ASDP, and Frick et al. (1992) found similar evidence for the role of parental ASDP in the association between poor parenting and offspring CD. Thus, it is possible that the association between family variables and CD may reflect a genetic rather than an environmental etiology (Lyton, 1990). Recent evidence from twin and adoption studies that have shown that many of the family risk factors for CD such as parenting behaviors and divorce are, in part, genetically inherited (McGue & Lykken, 1992; Plomin, 1995) also supports the hypothesis that CD may be genetically inherited.

Childhood antisocial behaviors have been shown to aggregate in families, as evidenced by significant sibling correlations (Rowe, Rodgers, & Meseck-Bushey, 1992; Szatmari, Boyle, & Offord, 1993) and associations between parental ASDP and offspring CD (Frick et al., 1992; Lahay, Piacentini, et al., 1988). The evidence from twin, adoption, and half-sibling studies concerning the relative magnitude of genetic versus shared family
environmental influences in the etiology of CD is mixed. There
depend on sex and age at diagnosis, but also that there are many
commonalities.

It is unclear to what extent methodological differences, such
differences in sample ascertainment (e.g., court-referred,
clinic-referred, or community volunteers), the age distribution of
the sample (e.g., preadolescent, adolescent, adult), the
methods of assessment (e.g., questionnaire, interview, official
record), or the source of information (e.g., self-report, parent
report) may explain differences in results of behavioral genetic
studies of childhood antisocial behaviors. In addition, most re-
cent twin studies of childhood antisocial behaviors (e.g., Edel-
brock, Rende, Plomin, & Thompson, 1995; Grove et al., 1990;
Lyons et al., 1995; Rowe, 1983) have used continuous measures
such as behavior or symptom counts rather than CD diagnoses.
The inheritance of antisocial or delinquent symptoms may or
may not be the same as the inheritance of the more extreme
pattern of behavior that corresponds to diagnosable CD. In other
words, whether CD is a discrete entity or an extreme of normally
occurring behavior remains to be determined (Hinshaw, 1994;
Hinshaw, Labey, & Hart, 1993). Behavioral genetic studies are
particularly valuable in addressing such questions. For example,
Nichols (1984) established that the inheritance of certain forms
of mental retardation differs from the inheritance of general
intelligence by examining the IQ of siblings of mildly versus severely retarded children (see Plomin, DeFries, & Mc
cLean, 1990). Similar methods can be used to determine the
extent to which continuous measures of antisocial and delin-
quent behaviors from the general population share the same risk
factors as CD.

One of the most consistent, but as yet unexplained, findings
for CD is the marked sex difference in the rate of the disorder.
This prevalence difference may reflect sex differences in the
etiology of CD. There is evidence to suggest that the risk factors
for CD may differ in boys and girls, but also that there are many
commonalities. Cloninger, Christiansen, Reich, and Gottesman
(1978) illustrated how family data can be used to examine sex
differences in the causes of psychiatric disorders by comparing
the similarity of male and female same-sex and unlike-sex (i.e.,
male-male, female-female, male-female) pairs of relatives.
They found that the familial causes of adult ASPD are largely
overlapping in men and women, but that women require more
familial risk factors before they are affected. Few behavioral
genetic studies have been capable of examining sex differences
in the etiology of childhood antisocial behaviors, either because
data on female probands (e.g., Lyons et al., 1995; Twito &
Stewart, 1982) or unlike-sex sibling or twin pairs (e.g., Edel-
brock et al., 1995; Rowe, 1983; Stevenson & Graham, 1988)
were not available. Some studies suggest that genetic influences
may account for a larger proportion of the variation in childhood
antisocial behaviors in boys than in girls (Silberg et al., 1994;
Stevenson & Graham, 1988; van den Oord, Boomsma, & Ver-
hulst, 1994), although this is not universally found (Rowe,
1983) and is inconsistent with the literature on childhood antisoc-
ial behavior in boys (including the Vietnam-era twin study) that
suggests very modest genetic influences. In sum, the general
question of sex differences in the etiology of CD remains unre-
solved because data on the development of CD in girls are
scant (Goodman & Kohlslorf, 1994; Rutter & Giller, 1983;
Zoccolillo, 1993).

The present study is an analysis of twin similarity for retro-
spectively assessed childhood CD in a sample of 2,682 male,
female, and unlike-sex adult Australian twin pairs. We examined
the following three main issues in the etiology of CD: (a) the
relative role of genes, shared family environment, and nonshared
individual-specific environment in the development of CD; (b)
the extent to which the etiology of CD is similar in boys and
girls; and (c) whether normal-range conduct problems or sub-
clinical manifestations of CD lie on the same liability dimension
and share the same risk factors as diagnosable CD.

Method

Participants

The participants were from the Australian National Health and Medical
Research Council Twin Register (ATR), a volunteer twin panel recruited
through the media, schools, and a variety of other sources (Jardine &
Martin, 1984). Three major health and behavior-related surveys have
been conducted to date with the ATR sample: a mailed questionnaire
survey in 1980–1981 (N = 8,183 individuals; 69% overall individual
response rate for questionnaire; Jardine & Martin, 1984), a follow-up
mailed questionnaire survey in 1988–1989 of all complete twin pairs
from the 1980–1981 survey (N = 6,327 individuals; 83% overall indi-
vidual response rate for questionnaire; Heath, Cloninger, & Martin,
1994; Heath & Martin, 1994), and a telephone interview survey conducted
in 1992–1993 (N = 5,889 individuals; 86% overall individual response
rate for interview; Heath, Bucholz, et al., 1994). Data obtained from
the telephone interview survey and the 1988–1989 questionnaire survey
were analyzed for the present study.

Telephone interviews were attempted with any twin pairs in which at
least one twin responded to the 1988–1989 mailed questionnaire
survey (N = 6,664 individuals; 86% overall individual response rate for
interview) or who had participated in an earlier alcohol challenge study
conducted in 1978–1979 (N = 412 individuals; 85% overall individual
response rate for interview; Heath and Martin, 1992; Martin, Oakeshott,
et al., 1985; Martins, Perl, et al., 1985). Telephone interviews were
completed with 5,889 individuals: 1,2041 men (mean age = 42.7 years, range = 28–89) and 3,846 women (mean age = 44.5 years, range = 27–90).

There were 2,685 complete twin pairs. Excluding three pairs in which one twin's CD diagnostic data were incomplete, there were 2,682 twin pairs available for analyses of twin similarity for CD (930 female monozygotic (MZ) twin pairs, 396 male MZ twin pairs, 533 female-female dizygotic (DZ) twin pairs, 231 male–male DZ twin pairs, and 592 female–male DZ twin pairs).

Preliminary data from an ongoing interview follow-up survey of a subsample of 604 individuals (326 men, 278 women) from twin pairs in which at least one twin had a lifetime history of the third revised Diagnostic and Statistical Manual of Mental Disorders (DSM–III–R) (American Psychiatric Association, 1987) alcohol dependence (assessed at the 1992–1993 interview) were analyzed for the purpose of evaluating the stability of the diagnoses of interest in the present study. The mean age at the 1992–1993 interview for this re-interviewed subsample was 39.4 years (range = 28–73).

Because the ATR is a volunteer twin register, it is possible that twins with a history of CD (who would be at heightened risk for adult antisocial behaviors) might have been undersampled. Possible sampling biases were examined by (a) comparing demographic and personality characteristics of the baseline sample (participants in the 1980–1981 questionnaire survey) to the interviewed sample, (b) comparing demographic and personality characteristics of twins whose cotwins did not participate in the baseline survey to twins who were concordant for participation in the baseline survey, and (c) comparing rates of CD in twins whose cotwin did not participate in the interview to the rate in twins who were concordant for participation in the interview.

The only demographic or personality characteristics on which the interviewed sample differed by more than 2% from the baseline sample were the proportion of MZ twins (men, 40% at baseline vs. 43% at interview; women, 49% at baseline vs. 51% at interview), the proportion of respondents born before 1930 (men, 13% at baseline vs. 9% at interview; women, 15% at baseline vs. 12% at interview), and the proportion of men with less than 11 years of education (20% at baseline vs. 16% at interview). Differences between the baseline and interviewed samples in marital status, religious affiliation, church attendance, and revised Eysenck Personality Questionnaire (EPQ-R; Eysenck, Eysenck, & Barrett, 1985) scale scores were modest.

Another check on sampling bias compared baseline demographic and personality characteristics in twins whose cotwin did not participate at baseline with twins whose whose participated at baseline. Assuming that twins are correlated for the baseline demographic and personality characteristics, differences between those who did and did not participate at baseline will be reflected in differences between cotwins of participants and cotwins of nonparticipants. The most consistent predictors of participation at both baseline and interview were being an MZ twin (in both men and women) and having a religious affiliation of “other Protestant” (in women only). There were no consistent associations of participation across surveys with age, education, marital status, church attendance, and personality characteristics.

Similarly, we examined the rates of CD among twins whose cotwin did not participate in the interview and among twins whose cotwin participated in the interview. If twins with CD were systematically undersampled, higher rates of CD would be expected among twins whose cotwin did not participate in the interview than among twins concordant for participation in the interview (assuming that CD status is correlated in twin pairs). There were 519 twin pairs who were discordant for participation in the telephone interview. The rates of CD among these unpaired twins did not differ significantly from the rates among the twins concordant for participation, \( \chi^2(1, N = 354–1,970) < 1.22, p > .27 \) for all six sex by zygosity groups. The combined CD rates across zygosity groups were 18.4% in concordant men versus 17.4% in discordant men, \( \chi^2(1, N = 2,041) = 0.11, p = .74 \), and 2.6% in concordant women versus 2.8% in discordant women, \( \chi^2(1, N = 3,842) = 0.02, p = .90 \).

Heath et al. (1996) examined the impact of cooperation bias and attrition on rates of CD and estimates of genetic and environmental influences on CD liability by weighting the interview data back to the characteristics of the baseline sample and to the distribution of educational attainment found in the general population of Australia. The rates of CD in the weighted data were very similar to the unweighted rates. Overall, the evidence suggests that this sample was not biased with respect to CD, although it is likely that severe cases of CD may have been underrepresented in this community-based sample. More important, twin correlations were nearly identical with the weighted and unweighted CD data. The analysis of Heath et al. (1996) suggests that the results of the present study cannot be dismissed solely on the basis of sampling bias.

Australia was the site of penal colonies for more than 168,000 British convicts from 1788–1868 (Wheeler, 1989). Despite this, the current crime rates in Australia are not particularly high compared with Europe, the United States, and Canada (Fingerhut & Kleiman, 1990; Mukherjee, 1996). In particular, the rates of violent crime such as homicide and robbery are much lower in Australia than in the United States and similar to the rates in England and Canada (Fingerhut & Kleiman, 1990; Mukherjee, 1996). The rates of nonviolent crimes such as burglary and larceny are roughly similar in Australia, the United States, England, and Canada (Mukherjee, 1996).

**Measures**

Measures used in the present study were obtained from two surveys of the ATR. CD was assessed in the telephone interview conducted in 1992–1993, and the measures of similarity of childhood experiences and frequency of contact with twin siblings as adults were assessed in the mailed questionnaire survey conducted in 1988–1989.

**Zygosity**

Twin zygosity diagnoses were based on questionnaire responses concerning physical similarity and how often the twins were mistaken for each other as children. This method of zygosity diagnosis has been shown to be about 95% accurate, as validated against blood-typing (Eaves, Eysenck, & Martin, 1989).

**CD**

The Semi-Structured Assessment for the Genetics of Alcoholism interview (SSAGA; Bucholz et al., 1994a), originally developed for the Collaborative Study on the Genetics of Alcoholism (COGA), was modified for use as a telephone interview in Australia. Interviews were administered by trained lay interviewers who were blind to the psychiatric status of the cotwin. Interviewers were supervised by a qualified clinical psychologist with 4 years of experience who reviewed all interview protocols. In addition, all interviews were tape-recorded and random interview tapes were reviewed for quality control.

Individual CD symptoms were assessed by telephone interview with the modified SSAGA. Participants were asked whether each of the 13 DSM–III–R CD symptoms ever occurred before the age of 18. Individuals who reported that they had the symptom before the age of 18 were asked whether it ever occurred before age 15, and those with 5 or more

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1. The reported sample size was 5,995 in previous articles. Subsequently, we discovered that 106 individuals (36 twin pairs and 34 single twins) were interviewed who did not meet the criteria for inclusion in the 1992–1993 interview survey. The 106 participants had responded to a mailed questionnaire of the ATR in 1980–1981 but had not participated (nor had their twin participated) in either the alcohol challenge study or the 1988–1989 mailed questionnaire survey. These 106 participants were excluded from the present study. However, when analyses were carried out with the additional participants included, the results did not change from those reported here.
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symptoms before age 15 were asked whether they ever had a 6-month period within which at least 3 symptoms co-occurred and persisted. The most common CD symptoms in this community-based sample were as follows: stolen without confrontation on more than one occasion (18%), often truant from school (15%), has deliberately engaged in setting fires (9%), often lies (6%), and has deliberately destroyed other’s property (6%). Symptoms that were very uncommon in this sample were as follows: stolen with confrontation of a victim (0.1%), has forced someone into sexual activity (0.1%), and has run away from home overnight at least twice (0.7%).

The individual CD symptoms were aggregated into lifetime CD diagnoses by computer algorithm. Three different CD diagnoses, ranging from a relatively broad to a relatively narrow definition of CD were examined. The focus of the present article was primarily on the broadest definition, which was endorsement of three or more CD symptoms ever occurring prior to age 18. This definition corresponds to the DSM-III-R CD diagnosis except for the 6-month symptom-duration criterion. The second definition of CD was endorsement of three or more CD symptoms ever occurring prior to age 15; this definition corresponds to the CD criterion for DSM-III-R ASPD. This operationalization of CD, albeit broad, is consistent with that used in the recent National Comorbidity Study (NCS; Kessler, Regier, & Nelson, 1995), an epidemiologic psychiatric interview survey of a national probability sample of 15–54-year-old men and women in the United States, and similar to the CD definition used in the Epidemiologic Catchment Area Survey (Robins & Regier, 1991). The third definition of CD was endorsement of three or more CD symptoms occurring prior to age 15 that co-occurred within and persisted for a period of at least 6 months; this definition may correspond to a syndrome more similar to those observed in treatment settings.

The 1-week interrater test–retest reliability (Yule’s Y, Spitznagel & Helzer, 1985) of CD with onset prior to age 15 assessed by the original SSAGA interview was .64–.65 (unpublished data from Bucholz et al., 1994), which is considered “fair” (Fleiss, 1981). Correlations in liability (see below) for this same definition of CD were .80–.82 (unpublished data from Bucholz et al., 1994). Stability of the retrospective reports of childhood CD was assessed in 604 participants from the present study who were re-interviewed on average 15 months later (range = 2–24 months). The stability of CD with onset prior to age 15, as indexed by Yule’s Y and the correlation in liability, was .76 and .83 and for CD with onset prior to age 18 was .67 and .78. The 15-month stability of the retrospective reports of childhood CD assessed in these adult participants was as high as the short-term reliabilities obtained for the SSAGA interview.

The use of retrospective assessments of CD in adults represents one of the main limitations of the present study. In fact, any study that assesses ASPD in adults, including the Epidemiologic Catchment Area (Robins & Regier, 1991) Study and NCS (Kessler et al., 1994), by design faces this limitation because the diagnosis of ASPD requires the diagnosis of childhood CD. Nonetheless, the reliability of such reports is acceptable, and there is evidence to suggest that concurrent (Hindelang, Hirschi, & Weiss, 1981; Hirschi, Hindelang, & Weis, 1980) and retrospective reports (Henry, Moffitt, Caspi, Langley, & Silva, 1994; Krueger et al., 1994) of delinquent behaviors are also reasonably valid. For example, 73% of 18-year-olds with a police record of arrest reported at interview that they had been arrested (Henry et al., 1994). Self-reported history of delinquent activity before age 15 correlated .43 with self-reported police contacts before age 15 retrospectively reported in men 36–55 years of age (Lyons, 1996), compared with a nearly equal correlation of .42 between self-reports of current delinquent activity and a past record of police contacts in 18-year-old men (Krueger et al., 1994). Self-reports of delinquent activity correspond well with official records of delinquent behaviors such as arrests or police contacts.

Similarity of experiences. In the 1988–1989 mailed questionnaire survey, twins were asked how often they shared the same room, had the same playmates, were dressed alike, and were in the same classes at school as children, with the response options of never, sometimes, usually, and always. Twin pairs who responded never or sometimes versus those who responded usually or always were compared in analyses. When twin responses conflicted (i.e., one twin’s response was never or sometimes and the other twin’s response was usually or always, the twin pair was placed in the never or sometimes category. In addition, twins were asked how often they currently saw or contacted their twin, and two dichotomous variables were created from the responses. Twins who currently saw each other at least once a week were contrasted with those who saw each other less frequently. Similarly, twins who currently contacted each other by telephone or letter at least once a week were contrasted with those who were in less frequent current contact.

Data Analysis

Two indexes of twin similarity for CD were computed: the probandwise concordance and the tetrachoric correlation. The probandwise concordance is an estimate of the probability that an individual has a CD diagnosis given that their cotwin has the diagnosis and is computed with only twin pairs with at least one affected member. The tetrachoric correlation, or correlation in liability (Olsson, 1979), is computed from the full two-by-two contingency table cross-classifying the CD status of a twin by the CD status of the cotwin. The tetrachoric correlation is used when one assumes that scores on the observed ordinal variables can be explained by underlying normally distributed variables (Falco1ner, 1965; Gottesman & Shields, 1967; Reich, Cloninger, & Guze, 1975).

In this case, the continuum underlying CD diagnostic status was presumed to be the liability to develop CD; participants were assigned a positive or negative CD diagnosis on the basis of whether or not their CD liability exceeded some arbitrary criterion or threshold (see Figure 1a). The tetrachoric correlation is the correlation between the presumed normally distributed latent CD liability between twin pairs.

In addition to DSM-III-R CD and the childhood criterion for DSM-III-R ASPD, we carried out twin analyses for a multiple-threshold model (Reich, James, & Morris, 1972) of CD liability (see Figure 1b). To test whether subclinical manifestations of CD (i.e., one or two CD symptoms endorsed) lie on the same liability dimension as the diagnosable disorder, the polythoracic correlation (the generalization of the tetrachoric correlation coefficient for polychotomous variables of three or more categories) between the number of CD symptoms (i.e., zero, one, two, or three or more symptoms) within twin pairs was calculated from the 4 x 4 contingency tables using PRELIS (Joreskog & Sorbom, 1988).

The assumption of an underlying distribution of liability was tested by comparing the observed frequencies in the 4 x 4 contingency tables with the frequencies expected assuming bivariate normal distributions of the underlying liability (Neale & Cardon, 1992). The results of these chi-square tests are presented, although they should not be overinterpreted because the statistical power to detect deviations from bivariate normality in ordinal data from a general population sample is low (Neale, Eaves, & Kendler, 1994).

Genetic and environmental structural equation models were fitted directly to the five twin contingency tables by maximum likelihood using the Mx program (Neale, 1995) to determine the relative roles of additive genetic (A), shared environmental (C; i.e., between family or common environment), and nonshared environmental (E; i.e., within family or individual-specific environment including measurement error) factors as sources of variation for CD liability. The fit of a series of nested submodels was compared by likelihood ratio chi-square to a most general full model, which allowed for additive genetic, shared environmental, and nonshared environmental factors as sources of variation (ACE model). For example, models in which the genetic influences (or shared environmental influences) were fixed at zero were compared with the model
containing all three sources of variation. When dropping parameters from a model did not result in a statistically significant decrement in model fit (i.e., there was a nonsignificant likelihood ratio chi-square test), then the reduced model was considered to give a better (i.e., more parsimonious) fit to the data. The final best fit model was the simplest model that was consistent with the data (see Heath et al., 1989; Neale & Cardon, 1992, for further details about model fitting). The overall adequacy of the fit of a particular model to the data was assessed by the goodness-of-fit chi-square test. Unlike most significance testing in psychological research, the goal is to obtain a nonsignificant goodness-of-fit chi-square; this indicates that the model cannot be rejected because it provides an adequate explanation of the covariance structure in the data. Genetic and environmental model fitting was carried out for DSM–III–R CD diagnoses and for a multiple-threshold model of CD liability.

Model fitting with data that included both male and female and unlike-sex twin pairs permitted tests of several hypotheses about possible sex differences in the etiology of CD. First, we tested whether the magnitude of genetic and environmental influences on CD liability differed in men and women. This was evaluated by comparing the fit of a model that allowed genetic and environmental parameter estimates to vary between men and women with the fit of a model that constrained genetic and environmental parameter estimates to be equal in the two sexes. Second, we tested whether the genetic and shared environmental mechanisms that influence the development of CD differed in men and women by testing whether the genetic correlation in unlike-sex twin pairs (r_g) was significantly different than the .50 that is expected for traits in which there is no sex limitation of genetic influences (while fixing the shared environmental correlation at .1) and, similarly, testing whether the shared environmental correlation in unlike-sex twin pairs (r_e) was significantly different from the .1 that is assumed for traits in which there are no sex-specific shared environmental influences (while fixing the genetic correlation at .5). Note that either r_g or r_e can be freely estimated in these models, but it is not possible to freely estimate both parameters simultaneously with data from MZ and DZ twins.

Confidence intervals for parameter estimates of additive genetic, shared environmental, and nonshared environmental effects were computed in Mx by determining the upper and lower bounds of a parameter that produced a statistically significant deterioration in model fit. This was done by fixing a parameter and comparing, by the likelihood ratio chi-square test, the fit of the model with the fixed parameter to the fit of the model in which the parameter was freely estimated. In the present article, 95% confidence limits of parameter estimates are presented for both the full and best-fit models. For both the full and best-fit models, confidence limits were computed within a saturated model that included all three sources of variation (additive genetic, shared environmental, and nonshared environmental influences). Confidence limits were computed separately for men and women for parameter estimates obtained under the full model, for best-fit models, confidence limits of parameter estimates were computed separately for men and women only if there was evidence for sex differences in the parameter estimate, otherwise confidence intervals were computed jointly for men and women.

This method of computing confidence intervals is preferred to comput-
ing the confidence interval as 1.96 times the standard error of the parameter estimate, because the standard error method assumes that the distribution of the parameter estimate is normal. In model fitting of twin data, the distribution of the parameter estimates are often skewed, resulting in confidence intervals that are asymmetrical around the parameter estimates.

MZ twins typically have more similar experiences than DZ twins; for example, they more frequently share playmates as children and are in more frequent contact as adults (Kendler, 1993). If such experiences are relevant to the development of CD, then the equal environmental similarity assumption of twin studies may not be correct for this disorder (Kendler, 1993; Kendler, Neale, Kessler, Heath, & Eaves, 1993). To test the equal environmental similarity assumption, logistic regression analyses predicting twin concordance for CD from similarity of childhood experiences were performed, controlling for zyosity, sex, and age. Because the assessment of CD was based on retrospective reports of the twins as adults, we also examined whether frequency of contact as adults predicted twin concordance for CD in similar logistic regression analyses.

Results

Lifetime Prevalence of CD

The prevalence of a self-reported childhood history of DSM– III–R CD was 18% in men and 3% in women. The sex difference in the rate of CD was highly significant, $\chi^2(1, N = 5,883) = 440.5, p < .001$. History of DSM–III–R CD also varied by birth cohort in both men, $\chi^2(4, N = 2,041) = 34.7, p < .001$, and women, $\chi^2(4, N = 3,842) = 37.3, p < .001$, with rates ranging from 22% and 5% in men and women under age 35, to 7% and 0.2% in men and women aged 60 years and older. The lifetime prevalence of CD with onset prior to age 15 was 13% in men and 2% in women, and when the requirement of a 6-month period of symptom clustering was imposed, the rates of CD fell to 4% in men and 0.6% in women. There were too few cases of CD with onset prior to age 15 and a 6-month period of symptom clustering to carry out analyses of twin similarity or genetic and environmental model fitting for this definition of CD, so for the remainder of this article only the two broader definitions of CD are used. We chose to focus primarily on the broadest definition of DSM–III–R CD because of the advantage of increased statistical power and doubts about participants’ ability to recall accurately information about the age of onset of symptoms over such long time intervals (on average a minimum of 25 years).

The rates of CD for Australian men and women in the present study were somewhat lower than rates reported for the general population of the United States in the NCS. Cottler, Kessler, and Nelson (1995) reported lifetime prevalence estimates of the childhood criterion for DSM–III–R ASPD (i.e., CD with onset prior to age 15) of 20% in men and 6% in women, compared with rates of 13% and 2% for the comparable definition of CD in this Australian sample. However, the older average age of this sample, compared with the NCS sample, may have contributed to the differences in CD rates. Once age was taken into consideration, the rate of CD in this Australian sample did not consistently differ with rates estimated for the general population of the United States. The rates of CD in the NCS study were 18% in men aged 35–44 years and 13% in men aged 45–54 years; in the Australian sample, the rates of CD before age 15 for men in these age groups were 15% and 14%. The rates of CD in the NCS study were 3% in women aged 35–44 years and 3% in women aged 45–54 years; in the Australian sample, the rates of CD before age 15 for women in these age groups were 2% and 2%. Only the rate in women 35–44 years of age was significantly different in the present sample compared with the rate in the NCS.

Twin Similarity for CD

Probands' concordances and correlations in liability for DSM–III–R CD are presented in Table 1. The MZ concordance was higher than the DZ concordance in both men (.53 vs .37) and women (.30 vs .18), but the difference attained statistical significance in men only, men: $\chi^2(1) = 6.2, p = .01$; women: $\chi^2(1) = 1.4, p = .24$. Similarly, the correlations in liability for DSM–III–R CD were higher among MZ than among DZ twins, but significantly so only among men, men: $.70 vs .37, \chi^2(1) = 6.7, p = .01$; women: $.68 vs .48, \chi^2(1) = 1.2, p = .28$. Correlations in liability for CD with an onset prior to age 15 were all within a standard error of the correlations in liability for DSM–III–R CD and had a similar pattern of MZ–DZ differences.

Sex Differences

Examination of the tetrachoric correlations in Table 1 suggests that the magnitude of genetic and environmental influences for CD do not differ in boys and girls. The tetrachoric correlations for DSM–III–R CD in male versus female MZ twin pairs did not differ significantly, .70 vs .68, $\chi^2(1) = 0.03, p = .87$, nor did the correlations in the same-sex male, female, and unlike-sex DZ twin pairs, .37 vs .48 vs .34, $\chi^2(2) = 0.44, p = .80$. In addition, the ratio of the unlike-sex DZ twin correlation to the geometric mean of the same-sex DZ correlations (see Cloninger et al., 1978) indicates that most of the familial influences relevant to the development of CD (81%) are shared by boys and girls. More rigorous tests of sex differences in the etiology of CD with model-fitting methods are presented below.

Fit of the Multiple-Threshold Model

Figure 1b presents the observed prevalences of each of the categories (zero, one, two, or three more conduct problems) used for the multiple-threshold model of CD for men and women, and the correlations in liability for the multiple-threshold model are presented in Table 1. The twin correlations for the multiple-threshold model were slightly reduced compared with the correlations in liability for DSM–III–R CD. In all five zyosity groups, the assumption of a bivariate normal liability distribution underlying the four ordered CD categories could not be rejected by a goodness-of-fit chi-square test.

Genetic and Environmental Model Fitting:

DSM–III–R CD

A full model allowing for additive genetic, shared environmental, and nonshared environmental influences for DSM–III–R CD fit the data well, goodness-of-fit $\chi^2(4) = 5.51, p = .24$, and yielded heritability estimates of 65% in men and 43% in
women and estimates of shared environmental influences of 4% in men and 25% in women (see Table 2). Setting the prevalences (i.e., threshold values) for CD to be equal in MZ male, DZ male, and males from unlike-sex pairs and, similarly, MZ female, DZ female, and females from unlike-sex pairs did not result in a significant deterioration in model fit, \( \chi^2(4) = 4.71, p = .32 \), and therefore, in all subsequent models, thresholds were constrained to be equal across zygosity groups. As expected, setting the prevalences for CD to be equal in men and women resulted in a substantial deterioration in model fit, \( \chi^2(1) = 308.80, p < .001 \); therefore, separate threshold values were estimated for men and women in all subsequent models.

When we fitted a full model allowing for sex differences in the etiology of DSM-III-R CD, estimates of either the between-sex genetic correlation \( (r_g) \) or the shared environmental correlation \( (r_c) \) were .44 and .56, respectively. Nonetheless, there was a not a significant deterioration in model fit when the estimate of \( r_g \) was fixed at .5, \( \chi^2(1) = 0.01, p = .93 \) or the estimate of \( r_c \) was fixed at 1, \( \chi^2(1) = 0.01, p = .93 \), and so in subsequent models \( r_g \) and \( r_c \) were fixed at these values. Setting equal the estimates of additive genetic, shared environmental, and non-shared environmental influences for men and women also did not result in a significant deterioration in model fit, \( \chi^2(2) = 0.37, p = .83 \); subsequent models constrained these estimates to be equivalent across sex. In sum, although inspection of the twin correlations suggested that there might be sex differences in the magnitude of genetic and environmental influences and in the specific additive genetic and shared environmental mechanisms that are important for the development of DSM-III-R CD, these sex differences were not statistically significant.

The fit of reduced models that accounted for DSM-III-R CD liability by the action of nonshared environmental influences only \( (E) \), shared and nonshared environmental influences \( (CE) \), and additive genetic and nonshared environmental influences \( (AE) \) were compared with the fit of a full model that included all three sources of variation \( (ACE) \). The E model, goodness-of-fit \( \chi^2(13) = 118.90, p < .001 \), and the CE model, goodness-of-fit \( \chi^2(12) = 23.98, p = .02 \), were both rejected by goodness-of-fit and likelihood ratio tests. In other words, genetic influences could not be dropped without a significant deterioration in model fit, \( \chi^2(1) = 13.38, p < .001 \). Eliminating shared environmental influences did not result in a significant deteriora-
tion in model fit, $\chi^2(1) = 0.05, p = .82$. A model that included additive genetic and nonshared environmental influences (AE) fit the data well, goodness-of-fit, $\chi^2(12) = 10.65, p = .56$, and was retained as the best-fitting model. The best-fitting model yielded variance estimates for additive genetic influences of 71% (95% confidence interval = 32–79%), for shared environmental influences of 0% (95% confidence interval = 0–32%), and for nonshared environmental influences of 29% (95% confidence interval = 21–41%).

Although not a primary focus in the present study, the effect of birth cohort was examined to determine whether it was an important source of variation for CD or a moderator of genetic and environmental influences on CD. When birth cohort was included in a full model, it accounted for 1% of the variance in CD liability in men (the estimates of the remaining variance components in the full model including birth cohort effects were as follows: A: 69%, C: 2%, and E: 28%) and 7% of the variance in women (A: 46%, C: 17%, and E: 30%). When the magnitude of genetic and environmental influences were estimated separately for different birth cohorts, the parameter estimates were not significantly different across cohorts, $\chi^2(8) = 12.03, p = .15$. In sum, birth cohort significantly contributed to the variation in liability for CD in the present sample but was not a significant moderator of genetic and environmental influences on risk for CD. Because the effect of birth cohort was relatively modest, models without birth cohort are reported in this article to simplify presentation. Note that the effect of birth cohort, when not separately estimated, will contribute to the estimate of shared environmental influences. Consequently, about one half of the estimated shared environmental influences for CD liability in men and about one third of the estimated shared environmental influences for CD liability in women in the full model in Table 2 can be attributed to birth cohort effects.

Model fitting for CD with onset prior to age 15 (the age corresponding to the childhood criteria for DSM-III-R ASPD) yielded very similar results as obtained for CD with onset prior to age 18, with the same estimates of additive genetic and nonshared environmental variances (71% and 29%, respectively) in the best-fitting model.

Genetic and Environmental Model Fitting: Multiple-Threshold Model of CD

The full and best-fitting models for the multiple-threshold model of CD liability are also presented in Table 2. The full ACE model fit the data well, goodness-of-fit $\chi^2(52) = 47.47, p = .65$, and yielded parameter estimates that were close to those for DSM-III-R CD liability, except that the estimate of additive genetic influences was somewhat reduced in men. The estimates of the between-sex genetic and environmental correlations were .46 and .80, respectively; these values were not significantly different from .5. $\chi^2(1) = 0.03, p = .87$, and 1, $\chi^2(1) = 0.03, p = .87$. Thus, there was little evidence to suggest that the specific genetic and environmental mechanisms associated with CD liability differed in men and women. However, a model that did not allow for sex differences in the estimates of the relative magnitude of additive genetic, shared environmental, and nonshared environmental influences (ACE model), goodness-of-fit $\chi^2(67) = 70.17, p = .37$, was rejected compared with a model that included sex-specific parameter estimates, $\chi^2(2) = 7.30, p = .03$.

We attempted to identify the most parsimonious sex-limitation model that could account for the data. A model that allowed for sex differences in parameter estimates but ruled out the influence of additive genetic factors (C, C, E, E model, where m = male and f = female), was rejected by both goodness-of-fit, $\chi^2(67) = 98.39, p = .01$, and likelihood ratio test. In other words, additive genetic influences could not be dropped from the model without a significant deterioration in fit, $\chi^2(2) = 35.52, p < .001$. Two parsimonious models fit the data without a significant deterioration in fit; an A, C, E, model, goodness-of-fit $\chi^2(67) = 64.86, p = .55$, and an A, C, E, model, goodness-of-fit $\chi^2(67) = 63.95, p = .58$. Because the fit of the latter model was slightly superior to the former, it is presented in Table 2 as the best-fitting model. The main difference in the parameter estimates between the two models was in the amount of variance apportioned to additive genetic and shared environmental influences in women; the parameter estimates for men were nearly identical for the two models (A: 53% vs. 52%, E: 47% vs. 48%). The best-fitting model for the multiple-threshold model of CD liability suggested that the magnitude of additive genetic influences were equal in men and women, accounting for 53% of the variance and that shared environmental influences were important only in women, accounting for 12% of the variance, with 35% of the variance in women accounted for by nonshared environmental influences. In the alternative model, additive genetic influences accounted for 66% and nonshared environmental influences accounted for 34% of the variation in CD liability in women.

Testing Assumptions of the Twin Method

The rate of DSM-III-R CD did not vary as a function of zygosity, women: $\chi^2(2) = 5.5, p = .06$; men: $\chi^2(2) = 2.2, p = .13$, which suggests that MZ and DZ twins do not differ in their risk for developing CD. Nonetheless, MZ twins reported more similar childhood experiences and more frequent contact as adults than DZ twins (e.g., 83% of female MZ twins usually or always shared the same playmates as children compared with 50% of like-sex female DZ twins). We examined whether this increased similarity of childhood experiences was associated with twin similarity for DSM-III-R CD. After controlling for zygosity, sex, and age, sharing the same room as children, $\chi^2(1) = 1.2, p = .27$, dressing alike as children, $\chi^2(1) = 0.94, p = .33$, and being in the same classes in school as children, $\chi^2(1) = 2.4, p = .12$, did not significantly predict twin concordance for DSM-III-R CD. There was a positive association, however, between sharing the same playmates as children, $\chi^2(1) = 5.2, p = .02$, and twin similarity for DSM-III-R CD. In addition,

2 The robustness of the best-fitting model choice was checked by repeating the model fitting with a different order to test for the significance of parameters. Thus, in addition to the procedure outlined in detail in the text, we also tested the significance of the genetic and shared environmental variance components prior to setting these parameter estimates equal in men and women. The conclusions and the best-fitting model remained the same regardless of the order of dropping nonsignificant parameters.
the frequency with which twins saw each other, \( \chi^2(1) = 4.6, p = .03 \), and contacted each other by telephone or letter, \( \chi^2(1) = 4.9, p = .03 \), as adults was also associated with twin similarity for DSM-III-R CD.

To shed more light on this issue, we examined the same-sex twin correlations stratified by similarity of experience. Among men, MZ twins who more often shared the same friends as children were more similar than MZ twins who less often shared the same friends, .76 versus .30, \( \chi^2(1) = 4.62, p = .03 \), but there was no difference in the similarity of DZ twins stratified by the frequency of sharing childhood peers, .46 versus .28, \( \chi^2(1) = 0.57, p = .45 \). When MZ–DZ comparisons conditional on sharing childhood peers were made, MZ twins who less often shared the same friends as children were no more similar than DZ twins who less often shared the same friends as children, .30 versus .28, \( \chi^2(1) = 0.00, p = .96 \), but there was a significant MZ–DZ difference among those twin pairs who more often shared the same friends as children, .76 versus .46, \( \chi^2(1) = 4.06, p = .04 \). Among male DZ twins, those who were more often in the same classes as children were more similar than those who were in the same classes less often, .72 versus .18, \( \chi^2(1) = 5.76, p = .02 \), but there were no differences among the MZ twin pairs, .72 versus .64, \( \chi^2(1) = 0.28, p = .60 \). MZ twins were more similar than DZ twins among those who were in the same classes less often, .64 versus .18, \( \chi^2(1) = 4.28, p = .04 \), but not among those who were more often in the same classes, .72 versus .72, \( \chi^2(1) = 0.00, p = .99 \). Among women, there were significant differences among MZ twins who differed in how often they dressed alike and were in the same classes as children, but these differences were opposite to what would be expected if greater similarity of experience leads to greater twin similarity for CD, that is, female twins who reported that they more often dressed alike or were in the same classes as children were less similar than those who reported that they less often dressed alike or were in the same classes.

In sum, these results suggest that, at least in men, the equal environmental similarity assumption may not be correct for twin studies of CD and that estimates of the genetic influences on CD should be interpreted with this caveat in mind. Although controlling for similarity of childhood experiences (namely, sharing the same friends and being in the same classes as children) effectively eliminated MZ–DZ twin differences in some subsamples, MZ–DZ differences remained significant in others. Even though genetic influences may be, to some extent, confounded with the greater environmental similarity of MZ than DZ twin pairs, genetic influences still are an important contributor to twin similarity for CD after controlling for the environmental similarity of twins. We discuss this issue further below.

Discussion

In a large community sample of adult Australian twins, we found evidence for a substantial genetic influence on the risk for DSM-III-R CD. The best-fitting model yielded a point estimate of heritability of 71%, with a 95% confidence interval of 32–79%. There was not a statistically significant effect of the shared environment in the best-fitting model of CD, but a modest effect of the shared environment on the risk for CD could not be excluded (95% confidence interval = 0–32%).

One point that should be emphasized is the lack of precision with which parameters such as heritability are estimated for psychiatric disorders (or any categorical trait with a prevalence substantially different from 50%), even with sample sizes as large as 2,682 twin pairs (Neale, Eaves, & Kendler, 1994). The heritability value itself should not be overly emphasized but rather the undeniable importance of genetic factors in the development of CD.

Although there was not a statistically significant effect of the shared environment in the development of DSM-III-R CD in the present study, shared environmental influences still may have accounted for up to 32% of the variation in liability. Twin studies can have limited power to detect the effect of the shared environment when it is estimated as a latent variable (i.e., inferred from the pattern of correlations in MZ and DZ twins) but can detect relatively modest effects of the shared environment when specific environmental risk factors are measured and included in the model fitting of twin data. For example, when childhood parental loss and separation was measured as a specific environmental risk factor, it accounted for a modest, but statistically significant, 2–5% of the variation in liability to anxiety disorders, depression (Kendler, Neale, Kessler, Heath, & Eaves, 1992), and alcoholism (Kendler et al., 1996) in women. Similarly, in the present study, there was a statistically significant effect of birth cohort, accounting for 1% and 7% of the variation in CD liability in men and women. Thus, although the overall effect of the shared environment was not statistically significant, the effect of a specific environmental risk factor significantly contributed to the variation in liability to develop CD. A substantial portion (30–50%) of the effect of the shared environment appeared to be due to differences in risk of CD for individuals growing up in different eras. With these cross-sectional data, it is unclear whether this effect represents true changes in the rates of CD in more recent eras or simply poorer recall of childhood misbehavior in older twins. Increased rates of self-reported CD in more recent birth cohorts in Australia are consistent with official statistics that document rises in crime (Biles, 1983; Mukherjee, 1996). Therefore, it is possible that sociocultural changes may be leading to increased risk for antisocial behaviors in youth.

The results of the present study agree with those of Rowe’s (1983) twin study of self-reported delinquency in mostly school-ascertained adolescents but conflict with a recent large twin study of CD retrospectively assessed in adult men (Lyons et al., 1995) that found a modest genetic effect and substantial shared environmental influences. The Vietnam-era twin panel sample, used in the Lyons et al. (1995) study, was identified from twin pairs in which both twins had been accepted for military service during the Vietnam era. Most participants in the Vietnam-era sample had completed high school, and only 9% met the childhood criteria for DSM-III-R ASPD (i.e., CD with onset prior to age 15), a much lower rate than is obtained in general population surveys of adults in the United States (Cottler, Kessler, & Nelson, 1995) and significantly lower than the rate of 14% of CD before age 15 in men from the same age range in the present study, \( \chi^2(1, N = 7,506) = 27.81, p < .001 \). The selected nature of the Vietnam-era twin panel may explain the markedly different results obtained in these studies. In addition, it is interesting to note that although there were not

Previous (nontwin) research has suggested that the causes of CD may differ in boys and girls (Goodman & Kohlsdorf, 1994; Rutter & Giller, 1983), and some twin studies suggest that the magnitude of genetic influences for CD may be greater in boys than in girls. In the present study, there was little evidence for sex differences in the etiology of CD. The magnitude of genetic and environmental influences for DSM–III–R CD did not vary significantly in men and women. In addition, the genetic and environmental mechanisms important for the development of CD appeared to be largely the same for both sexes.

In the present study, a multiple-threshold model of CD liability fit the data. This suggests that CD may not necessarily be a discrete entity but rather an extreme of the normal variation in conduct-disordered behavior found in the general population. The fit of this model suggests that the risk factors for and correlates and consequences of normal-range or subclinical forms of childhood antisocial behaviors are the same as those for CD. This is consistent with reports from the Epidemiologic Catchment Area Study (Robins & Regier, 1991), in which the number of childhood CD symptoms functioned as a continuum in predicting adult antisocial behaviors (Robins & Price, 1991; Robins, Tipp, & Przybeck, 1991), drug use (Robins & McEvoy, 1990), age of onset of alcohol and drug use (Robins & McEvoy, 1990), substance abuse (Robins & McEvoy, 1990), alcohol and drug abuse or dependence (Robins & Price, 1991), anxiety disorders (Robins & Price, 1991), somatization disorder (Robins & Price, 1991), depression (Robins & Price, 1991), and schizophrenia (Robins & Price, 1991). Results of studies of symptom counts of antisocial, delinquent, or externalizing behaviors can probably be generalized to CD. This alternative approach to assessing childhood antisocial behaviors has the advantage that it uses more of the available information in the data (i.e., the number of different behaviors rather than whether or not the child has exceeded a single diagnostic threshold) and, thus, has increased statistical power (Neale, Eaves, & Kendler, 1994). For example, in the current study, the 95% confidence intervals for parameter estimates under the multiple-threshold model were much narrower than under the DSM–III–R model.

Even though this study suggests that CD, for the most part, can be thought of as the extreme of a continuum of childhood antisocial behavior, this does not rule out the possibility of etiologically distinct forms of CD. Community-based studies may have limited power to detect etiologically distinct but relatively rare forms of a disorder as being discontinuous with more common forms of the disorder. For example, severe physical or sexual abuse or other trauma may cause CD in the absence of genetic vulnerability, or severe CD may be etiologically distinct from less severe forms of CD. Furthermore, although the multiple-threshold model of CD liability could not be rejected, we found that the correlation in MZ men was somewhat reduced for the multiple-threshold model compared with the DSM–III–R model of CD, resulting in a lower estimate of genetic influences for CD in men. This suggests the possibility that there may be unique genetic influences, in addition to those that contribute to normal-range variation in childhood antisocial behaviors, that predispose to diagnosable CD in boys.

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validity of twin method assumptions

The results of the present study must be evaluated with the assumptions of the twin method in mind. Although there is ample evidence for the validity of these assumptions for many psychological domains (Kendler, 1993; Plomin, DeFries, & McClearn, 1990), it is worthwhile to consider the assumptions of the twin method specifically with respect to childhood CD.

The first assumption of the twin method that must be considered is whether twins are representative of the general (nontwin) population with respect to CD or, more broadly, with respect to possible risk factors for CD. There is some evidence to suggest that there is a slight excess of CD or similar behavioral problems in twins compared with singletons, both in community and psychiatric samples (Gau, Silberg, Erickson, & Hewitt, 1992; Rutter & Redshaw, 1991; Simonoff, 1992), although van den Oord, Koot, Boomsma, Verhulst, and Orlebeke (1995) found lower levels of mother-rated oppositional behaviors in preschool twins compared with singletons. In addition, twins have slightly lower IQ scores than singletons on average and are more likely to have pre- and perinatal birth complications, language difficulties (Rutter & Redshaw, 1991; Simonoff, 1992), and depressed mothers (Thorpe, Golding, MacGillivray, & Greenwood, 1991). These findings are consistent with and may, in part, explain the slight excess of childhood antisocial behaviors in twins compared with singletons. The generalizability of twin studies of childhood psychopathology is still a matter open to debate (Rutter, Simonoff, & Silberg, 1993). It should be stated, however, that if twinning is a risk factor for CD, this would most likely lead to reduced estimates of the contribution of genetic factors and inflated estimates of either shared or nonshared environmental factors in its etiology. Thus, the violation of this twin-method assumption is unlikely to be of importance in the present study.

The other major assumption of the twin method that must be considered is that the environmental influences relevant to the development of CD are not more similar in MZ than DZ twin pairs. When this assumption is not met, estimates of genetic influences obtained from twin studies may be inflated. Indeed, in the present study, we found that MZ twins were more likely than DZ twins to have the same friends as children and that this significantly predicted twin concordance for CD. This is consistent with a large literature on the association between peer relationships and CD. The interpretation of this association, however, is somewhat problematic because the direction of causation is unclear. Although associating with delinquent peers may cause a child to engage in delinquent behaviors, there is also evidence in favor of the alternative hypothesis—that children associate with peers who are similar to them (Rowe, 1994). Delinquent children are more likely to associate with delinquent peers. Baker and Daniels (1990; see also Rowe, 1994), comparing MZ twins, DZ twins, full siblings, and adopted siblings for similarity of peer group characteristics, showed that more

3 We thank Mike Neale for this idea.
Implications

The results of the present study have important implications for the types of risk factors that are likely to contribute to the liability to develop CD. For example, we could not reject the possibility that associating with delinquent peers may increase a child’s risk of developing CD, although recent longitudinal studies of delinquent or disruptive behaviors in children suggest that the influence of peers is modest once the prior level of delinquent behavior in the child is taken into consideration (Hozza, Molina, Bukowski, & Sippola, 1995; Tremblay, Masse, Vitaco, & Dobkin, 1995). The relationship between peer influences and CD most likely reflects both genetic (assortative friendship) and environmental (peer imitation) mechanisms. Extended twin designs that also gather information from the friends of twins may help to better understand this important association.

Another promising area for future research is the identification of traits that mediate the genetic influences on CD. Personality traits such as deceitfulness (Rowe, 1986), anger or aggression (Krueger et al., 1994; Rowe, 1986), impulsivity (Krueger et al., 1994; Rowe, 1986; Tremblay, Pihl, Vitaco, & Dobkins 1994; White et al., 1994), and nonconformity (Krueger et al., 1994) have been associated with childhood antisocial behaviors, and Rowe (1986) demonstrated that the association of some of these traits with delinquent behaviors is, in part, genetically mediated. Goldsmith and Gottesman (1996) suggest several other promising candidate traits that may constitute the genetic diathesis underlying CD: need for stimulation, activity level, anxiety proneness, and frustration tolerance. These authors also suggest pathways by which genes might influence conduct-disordered behavior. Clearly, the route through which genes influence CD liability is complex.

This study suggests that the association of parent characteristics and parenting behaviors with CD in offspring may be due to genetic rather than shared family environmental mechanisms. If this is true, then modification of parent characteristics may not have the anticipated effect of reducing antisocial behaviors in their children. Future twin-family studies of CD that include specific family environmental risk factors (e.g., Kendler et al., 1996) will be helpful to further understand the causal significance of these putative risk factors in the development of CD.

Even if shared environmental influences play only a minor role in the development of CD in the majority of individuals, this should not lead to therapeutic nihilism (Mecheil, 1972, p. 10). The genetic etiology of a disorder does not imply immutability, and there are many examples of genetically influenced traits that have been modified through environmental interventions: Children’s smiles are improved with braces, their vision is improved with eyeglasses, their hearing is improved with hearing aids, and their overall health and growth is improved through proper nutrition. Even single-gene disorders are modifiable through environmental interventions: Babies born with phenylketonuria (PKU), galactosemia (see Plomin et al., 1990), and other metabolic disorders are often able to lead normal lives because of effective therapies. Effective treatments were developed only after the etiology of these disorders was known. Understanding the etiology of CD, and particularly the direction of causation of the association of CD with its putative risk factors, will very likely be helpful in the development of effective interventions for this disorder as well.

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