The Structure of Genetic and Environmental Risk Factors for Syndromal and Subsyndromal Common DSM-IV Axis I and All Axis II Disorders

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Objective: The authors sought to clarify the structure of the genetic and environmental risk factors for 22 DSM-IV disorders: 12 common axis I disorders and all 10 axis II disorders.

Method: The authors examined syndromal and subsyndromal axis I diagnoses and five categories reflecting number of endorsed criteria for axis II disorders in 2,111 personally interviewed young adult members of the Norwegian Institute of Public Health Twin Panel.

Results: Four correlated genetic factors were identified: axis I internalizing, axis II internalizing, axis I externalizing, and axis II externalizing. Factors 1 and 2 and factors 3 and 4 were moderately correlated, supporting the importance of the internalizing-externalizing distinction. Five disorders had substantial loadings on two factors: borderline personality disorder (factors 3 and 4), somatoform disorder (factors 1 and 2), paranoid and dependent personality disorders (factors 2 and 4), and eating disorders (factors 1 and 4). Three correlated environmental factors were identified: axis II disorders, axis I internalizing disorders, and externalizing disorders versus anxiety disorders.

Conclusions: Common axis I and II psychiatric disorders have a coherent underlying genetic structure that reflects two major dimensions: internalizing versus externalizing, and axis I versus axis II. The underlying structure of environmental influences is quite different. The organization of common psychiatric disorders into coherent groups results largely from genetic, not environmental, factors. These results should be interpreted in the context of unavoidable limitations of current statistical methods applied to this number of diagnostic categories.

Psyciatric disorders are clinical-historical constructs whose etiology and pathophysiology are largely unknown, and hence most psychiatric nosologies, including DSM-IV (1) and ICD-10 (2), arrange disorders into categories primarily on the basis of clinical similarities. Our field has long hoped for an etiologically based classification of psychiatric disorders. Of the possible organizing principles for such an approach, familial/genetic factors have frequently been emphasized (3–5).

The potential utility of examining the broad structure of psychiatric and substance use disorders was first demonstrated by Krueger and colleagues (6–8) and then by others (e.g., references 9, 10). These analyses, examining a range of epidemiological samples, provided consistent evidence that common psychiatric disorders could be divided into two broad categories: internalizing disorders—dominated by major depression and anxiety disorders—and externalizing disorders—dominated by antisocial personality disorder and drug and alcohol use disorders.

Such epidemiological investigations, while instructive, do not directly provide insight into the causes of the patterns of co-occurrence. By contrast, similar studies performed in genetically informative samples, such as twins, can distinguish between the structures of psychiatric disorders produced by genetic factors on the one hand and environmental factors on the other. However, it might be asked: What is the value of such an approach in the genomic era? Examining the impact of individual molecular variants on risk for multiple psychiatric disorders will explain only very small proportions of the shared genetic variance (11). Twin studies, by contrast, assess aggregate genetic effects and therefore examine the degree of sharing across disorders of all genetic risk variants. While molecular methods can lead to a clarification of common pathophysiological pathways, the global questions that psychiatric nosologists have traditionally been interested in (e.g., how closely related genetically are two disorders?) can be best addressed at the aggregate level using genetically informative designs like twin studies (12).

One previous study examined seven common psychiatric and substance use disorders in members of the Virginia Twin Study of Psychiatric and Substance Use Disorders (13). Consistent with previous epidemiological investigations, that study identified two genetic factors that loaded strongly on, respectively, three internalizing disorders (major depression, generalized anxiety disorder, and pho-
bias) and four externalizing disorders (alcohol dependence, drug abuse or dependence, antisocial personality disorder, and conduct disorder). A parallel analysis of the effects of the environmental factors did not produce clear internalizing and externalizing factors, which suggests that genetic rather than environmental factors are responsible for the coherent underlying structure of common psychiatric disorders. Other researchers have documented the genetic coherence of an externalizing dimension of psychopathology (e.g., references 14, 15).

The present study, conducted in the Norwegian Institute of Public Health Twin Panel (NIPHTP), represents a follow-up and expansion of our earlier effort (13) in two critical ways. First, we examined a much broader array of common axis I disorders. Second, while the previous study included only one axis II disorder (antisocial personality disorder), the present analysis includes all 10 of the DSM-IV personality disorders. This addition permitted us, for the first time, to study systematically the genetic and environmental relationships between axis I and axis II disorders, a key component of the current conceptual framework for psychiatric disorders first introduced in DSM-III (16). One limitation of our approach is worth highlighting. Despite our relatively large sample, many of the individual disorders we wished to study were too rare to be examined solely at a fully syndromal level. Therefore, for the personality disorders, as we have done in the past (17–19) and others have advocated (20–24), we examined the number of endorsed criteria. For most of the axis I disorders, we also examined patients with subsyndromal cases (and show that such individuals had disorders that reflected milder manifestations of the same underlying liability as those with fully syndromal disorders).

The goal of this study, then, was to investigate the underlying genetic and environmental structure of a large proportion of common axis I DSM-IV disorders and all DSM-IV axis II disorders. We sought to clarify for the first time the broad structure of common axis I and axis II psychiatric syndromal and subsyndromal disorders as seen from an etiological perspective, in this case from a genetic point of view.

Method

Sample and Assessment Methods

Twins were recruited from the NIPHTP (25). Twins in the NIPHTP were identified through the Norwegian National Medical Birth Registry, which was established on January 1, 1967, and receives mandatory notification of all live births. Questionnaire studies were previously conducted in 1992 (twins born 1967–1974) and in 1998 (twins born 1967–1979). Altogether, 12,698 twins received the second questionnaire, and 8,045 (3,334 pairs and 1,377 single responders) responded after one reminder (cooperation rate, 63%).

The data for this analysis are from an interview study, conducted from 1999 to 2004, assessing DSM-IV axis I and axis II disorders (26). Interviewers were largely senior clinical psychology students at the end of their 6-year training course (including at least 6 months of clinical practice) and psychiatric nurses with years of clinical experience. They were trained by professionals with extensive experience with the instrument. The interviews were mostly conducted face-to-face, although for practical reasons 231 interviews (8.3%) were conducted by telephone. Each twin in a pair was interviewed by a different interviewer.

As outlined in detail elsewhere (26), the 6,442 eligible participants were defined as the 3,153 complete pairs in which both members completed the second questionnaire and agreed to be contacted again, as well as 68 pairs unintentionally drawn directly from the NIPHTP.

Altogether, 2,794 twins (44% of those eligible) were interviewed. Noncooperation was overwhelmingly the result of nonresponse to the written invitation; active refusals were rare (0.8%) (26). The study was approved by the Norwegian Data Inspectorate and the Regional Ethical Committee, and written informed consent was obtained from all participants after they received a complete description of the study. Of those interviewed, 36.5% were male, and the mean age was 28.2 years (SD=3.9).

As outlined elsewhere (27), zygoity was determined by the use of questionnaire items for the entire sample (28) and by microsatellite markers for 676 of the like-sex pairs, which, when used together in a discriminant analysis for participants for whom DNA was unavailable, predicted a zygosity misclassification rate of -1% of pairs, a rate far too low to substantially bias results (29).

These analyses included only the same-sex pairs from this sample—2,111 individuals, including both members of 669 monzygotic and 377 dizygotic pairs and 19 individual twins without their co-twin. Only twin pairs in which both twins initially had agreed to participate were interviewed; for the 19 individual twins, the co-twin changed his or her mind after the initial consent.

Axis I Disorders

Axis I disorders other than conduct disorder were assessed using a Norwegian computerized version of the Munich-Composite International Diagnostic Interview (M-CIDI) (30)—a comprehensive structured diagnostic interview assessing DSM-IV axis I disorders (1) that has been shown to have good test-retest and interrater reliability (31–33). Both the paper-and-pencil version and the computerized version of the M-CIDI have previously been used in Norway (34, 35)

Twelve axis I disorders were included in these analyses: major depression, dysthymia, panic disorder, agoraphobia, specific phobia, social phobia, generalized anxiety disorder, eating disorders, somatoform disorder, alcohol abuse or dependence, illicit drug abuse or dependence, and conduct disorder. Except for conduct disorder, initial analyses of these disorders were performed using DSM-IV diagnoses automatically generated by the M-CIDI data program. However, stable solutions were unobtainable because of small or zero cell frequencies for a number of these diagnoses. We therefore created three ordered categories (unaffected, subsyndromal, and fully syndromal) for 10 of these disorders, which substantially improved the stability of our estimates. Our definitions for the subsyndromal categories are detailed briefly in Table 1 and in more detail in Table S1 in the data supplement that accompanies the online edition of this article.

We assessed the validity of our subsyndromal categories in two ways. First, we compared all the cross-disorder phenotypic correlations using the original dichotomous full-syndrome variables (tetrachoric correlations) with the three-category variables (polychoric correlations); 95% confidence intervals for these two correlations overlapped for 65 of 66 correlations. Second, using a multiple-threshold model in PRELIS 2.3 (36), we tested whether the three categories represented differing levels of severity on a single continuum of liability. This test failed at the 5% level three of 46 times, consistent with chance expectations (37).

The phenotypic tetrachoric and polychoric correlations for this sample between all 22 axis I and II disorders examined in these analyses are listed in Table S2 in the online data supplement.
Axis II Disorders

A Norwegian version of the Structured Interview for DSM-IV Personality (SIDP-IV) (38) was used to assess all 10 DSM-IV personality disorders and conduct disorder. The DSM-III-R and DSM-IV versions of this interview have been used previously in large-scale studies in Norway (39, 40). The SIDP-IV, a comprehensive semistructured diagnostic interview for the assessment of DSM-IV personality disorders, contains nonpejorative questions organized into topical sections rather than by individual personality disorder, thereby improving the interview flow. The SIDP-IV interview was conducted after the M-CIDI, which helped to distinguish long-standing behaviors from temporary states resulting from axis I disorders.

The SIDP-IV uses the “5-year rule,” meaning that behaviors, cognitions, and feelings that predominated for most of the past 5 years are judged to be representative of an individual’s personality. Each DSM-IV criterion is scored on a 4-point scale (0=absent, 1=subthreshold, 2=present, or 3=strongly present). To keep results parallel with other personality disorders, we examined only the A criterion for antisocial personality disorder.

With traditional cutoff scores, too few individuals met full DSM-IV criteria for the 10 personality disorders for statistical analysis (17–19). We therefore modeled the personality disorders as an ordinal count of the number of positively endorsed criteria. Furthermore, defining a criterion to be present with a score of 1 or higher produced more stable results than using a cutoff of 2 or higher. This approach is justified by results from previous studies of these 10 personality disorders (17–19) in which, using a multiple-threshold model, we showed that the four response options for scoring individual personality disorder criteria reflected varying levels of “severity” on a single continuum of liability.

Because few individuals endorsed most of the criteria for individual personality disorders, we collapsed the total criterion count into five categories to reduce the frequency of null cells. We have also tested the validity of this approach by examining the fit of the multiple-threshold model, which asks whether the number of endorsed criteria reflects differences of severity on a single normal continuum of liability. This assumption was supported for all 10 personality disorders (17–19). For ease of expression, we refer in this article to “personality disorders” in place of the more accurate but cumbersome term “five categories of endorsed criteria for personality disorders.” We previously reported the high interrater reliability for the assessed personality disorder obtained by two raters scoring 70 audiotaped interviews (27) (intraclass correlations for number of endorsed criteria ranged from 0.81 to 0.96).

Statistical Methods

Our analytic approach involved three major steps: 1) estimating polygenic correlations for 44 (2×22) variables, including within-twin cross-disorder, cross-twin-within-disorder, and cross-twin cross-disorder correlations, for monozygotic and dizygotic twins separately; 2) estimating genetic and environmental correlations between all 22 disorders, based on multivariate biometric modeling; and 3) applying exploratory factor analysis to the resulting genetic and environmental correlation matrices.

First, monozygotic and dizygotic polygenic correlations with corresponding asymptotic weights were estimated in Mplus 5.21 for the monozygotic and same-sex dizygotic twin pair data (41). The robust weighted least squares mean and variance estimator was used. Under this method, all twin correlations for all disorders variables are estimated pairwise using all available ordinal raw data for each combination of variables. The weights are the estimated variances of these correlation parameters. These were obtained in Mplus using the TECH3 output and savedata options. These asymptotic variances up- or down-weight the contribution of each of the respective polygenic correlations.

Next, a saturated Cholesky decomposition of the monozygotic and dizygotic twin correlations among our 22 disorders was performed in Mx. A diagonally weighted least squares fit function was implemented in Mx (42) to maximize the agreement between the observed statistics and those predicted by the model. The squared deviations between observed and expected correlations were weighted by the inverse of the asymptotic covariances of each statistic; these weights were computed using Mplus. Because of the large number of variables in the model, we had to use limited-information diagonally weighted least squares

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### TABLE 1. Summary Criteria for Subthreshold Forms of Common Axis I Disorders

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Summary Criteria for Subthreshold Form of the Disorder</th>
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</thead>
<tbody>
<tr>
<td>Panic disorder</td>
<td>Assessed using two probe questions evaluating criterion A1 and required the experience of a panic attack without being in danger.</td>
</tr>
<tr>
<td>Agoraphobia</td>
<td>Assessed by a probe question evaluating criterion A and required reporting a strong fear of being away from home, traveling in a bus or train, being in a public place, traveling alone, or crossing a bridge.</td>
</tr>
<tr>
<td>Phobia</td>
<td>Assessed by a probe question evaluating criterion A and required an unusually strong fear or avoidance of living things, the sight of blood, getting an injection, going to the dentist or hospital, heights, storms, thunder or lightning, being in water, flying in an airplane, being in a closed space, or any other specified situation.</td>
</tr>
<tr>
<td>Social phobia</td>
<td>Assessed by two probe questions for criterion A and required an unusually strong fear or avoidance of seven possible social situations (e.g., eating, drinking, or writing while being watched, participating or speaking in a meeting) in addition to a fear of being in nine possible ways (e.g., showing anxiety, throwing up, losing control of bladder or bowels, blushing).</td>
</tr>
<tr>
<td>Generalized anxiety disorder</td>
<td>Assessed by a probe question evaluating criterion A and required the occurrence of a period of feeling worried, tense, or anxious most of the time for a minimum duration of 3 months.</td>
</tr>
<tr>
<td>Eating disorder</td>
<td>Assessed from criterion A for anorexia nervosa and criteria A and B for bulimia nervosa and required either loss of at least 6.5 kg and being perceived as too thin, or a period of binge eating and strict dieting, a lot of exercise, use of medication, vomiting, or fasting.</td>
</tr>
<tr>
<td>Somatoform disorder</td>
<td>Assessed by probe questions addressing criterion A of conversion disorder, criterion 5 of dissociative disorder not otherwise specified, criterion B of somatization disorder, criterion A of undifferentiated somatization disorder, criterion A of hypochondriasis, and criterion A of pain disorder.</td>
</tr>
<tr>
<td>Alcohol abuse or dependence</td>
<td>Required the admission of having had periods of drinking too much and consuming at least five units of alcohol per day twice a month or more.</td>
</tr>
<tr>
<td>Drug abuse or dependence</td>
<td>Required the use of illicit or improper use of prescribed drugs 10 or more times to get high. Nicotine dependence is not included.</td>
</tr>
<tr>
<td>Conduct disorder</td>
<td>Rating by interviewer as subthreshold (≥1) conduct disorder A criterion.</td>
</tr>
</tbody>
</table>
### TABLE 2. Estimated Proportion of Variance in Liability to Common DSM-IV Axis I and All Axis II Disorders Due to Genetic Effects (Heritability) and Individual-Specific Environment

<table>
<thead>
<tr>
<th>Disorder</th>
<th>Heritability (a²)</th>
<th>Individual-Specific Environmental Effects (e²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Major depression</td>
<td>0.43</td>
<td>0.57</td>
</tr>
<tr>
<td>Generalized anxiety disorder</td>
<td>0.51</td>
<td>0.49</td>
</tr>
<tr>
<td>Eating disorder</td>
<td>0.42</td>
<td>0.58</td>
</tr>
<tr>
<td>Social phobia</td>
<td>0.57</td>
<td>0.43</td>
</tr>
<tr>
<td>Agoraphobia</td>
<td>0.60</td>
<td>0.40</td>
</tr>
<tr>
<td>Specific phobia</td>
<td>0.55</td>
<td>0.45</td>
</tr>
<tr>
<td>Somatoform disorder</td>
<td>0.44</td>
<td>0.56</td>
</tr>
<tr>
<td>Dysthymia</td>
<td>0.28</td>
<td>0.72</td>
</tr>
<tr>
<td>Panic disorder</td>
<td>0.55</td>
<td>0.45</td>
</tr>
<tr>
<td>Alcohol abuse or dependence</td>
<td>0.52</td>
<td>0.48</td>
</tr>
<tr>
<td>Drug abuse or dependence</td>
<td>0.59</td>
<td>0.41</td>
</tr>
<tr>
<td>Conduct disorder</td>
<td>0.57</td>
<td>0.43</td>
</tr>
<tr>
<td>Paranoid personality disorder</td>
<td>0.29</td>
<td>0.71</td>
</tr>
<tr>
<td>Schizoid personality disorder</td>
<td>0.34</td>
<td>0.66</td>
</tr>
<tr>
<td>Schizotypal personality disorder</td>
<td>0.38</td>
<td>0.62</td>
</tr>
<tr>
<td>Antisocial personality disorder</td>
<td>0.50</td>
<td>0.50</td>
</tr>
<tr>
<td>Borderline personality disorder</td>
<td>0.49</td>
<td>0.51</td>
</tr>
<tr>
<td>Histrionic personality disorder</td>
<td>0.32</td>
<td>0.68</td>
</tr>
<tr>
<td>Narcissistic personality disorder</td>
<td>0.35</td>
<td>0.65</td>
</tr>
<tr>
<td>Dependent personality disorder</td>
<td>0.37</td>
<td>0.63</td>
</tr>
<tr>
<td>Avoidant personality disorder</td>
<td>0.47</td>
<td>0.53</td>
</tr>
<tr>
<td>Obsessive-compulsive personality disorder</td>
<td>0.34</td>
<td>0.66</td>
</tr>
</tbody>
</table>

The estimated genetic and environmental correlation matrices are presented in Tables S3 and S4 in the online data supplement. Table 2 presents the heritability of the individual disorders estimated from our model.

#### Results

The estimated genetic and environmental correlation matrices are presented in Tables S3 and S4 in the online data supplement. Table 2 presents the heritability of the individual disorders estimated from our model.

### Genetic Factors

Parameter estimates from the four genetic factors are presented in Table 3 and in Figure 1 (which includes all paths accounting for ≥10% of the genetic variance, that is, with loadings >0.316). Three broad patterns of findings are noteworthy. First, the four factors are all coherent and substantively interpretable. Factor 1 has strong loadings on four anxiety disorders (panic disorder, agoraphobia, specific phobia, and generalized anxiety disorder), major depression, eating disorders, and somatoform disorders. This factor reflects axis I internalizing disorders. Factor 2 is dominated by four personality disorders—schizoid, schizotypal, avoidant, and dependent—as well as dysthymia and social phobia. We name this factor axis II internalizing disorders, although it also contains two disorders traditionally placed on axis I. Factor 3 has strong loadings on four disorders reflecting antisocial behaviors (conduct disorder and antisocial personality disorder) and substance use disorders (alcohol and drug abuse or dependence). This factor identifies an axis I externalizing disorders dimension although it includes one category (antisocial personality disorder) that is traditionally placed on axis II. Factor 4 contains the five remaining personality disorders—histrionic, narcissistic, obsessive-compulsive, borderline, and paranoid—and is best interpreted as an axis II externalizing disorders dimension.

Second, the interfactor correlations quantify the degree of sharing of genetic risk factors for these four groupings of disorders. Most closely related are the two internalizing disorder factors (correlation=0.49) followed by the two externalizing factors (0.38). The two axis II groups are moderately intercorrelated (0.36), and the correlation between the two axis I groups was quite low (0.16).

Third, five disorders had notable cross-loadings on other factors: somatoform disorder (loadings from both factors 1 and 2), borderline personality disorder (factors 3 and 4), eating disorders (factors 1 and 4), and both paranoid and dependent personality disorders (factors 2 and 4).

### Environmental Factors

The factor loadings for the three environmental factors are presented in Table 4 and Figure 2 (which includes all
Fourth, the six disorders that had substantial loadings on two factors provide further insights into the structure of environmental risks for psychiatric disorders. Antisocial personality disorder was transitional between factors 1 and 3, and agoraphobia and panic disorder between factors 2 and 3. Factor 3 had secondary negative cross-loadings on two additional anxiety/dysphoria disorders: social phobia and dependent personality disorder.

Discussion

Our goal in this study was to clarify the structure of genetic and environmental risk factors for syndromal and subsyndromal common axis I disorders and all axis II personality disorders as assessed by a criterion count. Using multivariate twin analyses, we identified, from the 22 disorders examined, four coherent genetic factors: axis I internalizing, axis II internalizing, axis I externalizing, and axis II externalizing.

Although not without important limitations (see below), these findings provide, for the first time, a view of the etiological structure of a substantial proportion of common psychiatric disorders. Furthermore, this structure, especially the genetic factors, is coherent and clinically sensible. The structure of the genetic risks for these disorders is neither extremely simple (e.g., just one dimension of underlying risk) nor bewilderingly complex. Of the many interesting results from these analyses, three are particularly noteworthy.
First, we replicated and extended the results of our earlier multivariate twin analysis, which included only seven disorders but clearly identified genetic internalizing and externalizing factors (13). The present study provides further support for the importance and generalizability of the internalizing and externalizing genetic dimensions of risk for common psychiatric disorders. While we identified separate internalizing and externalizing factors for axis I and II disorders, they were moderately intercorrelated.

Second, our results provide—for the first time to our knowledge—some support, from a genetic perspective, for the decision in DSM-III to distinguish between axis I and axis II disorders. The genetic substrate for axis II disorders is, in our analyses, at least partially separable from those factors that predispose to axis I disorders. However, the axis I and II disorders that loaded on our genetic factors are not isomorphic with those articulated by DSM-III. Two axis I disorders—dysthymia and social phobia—were included in the internalizing axis II cluster. The concept of dysthymia evolved in part from the concept of “depressive personality” (16, 43). Our results suggest that from a genetic perspective, it may be better placed with the personality disorders than in the mood disorders section. A debate has long simmered about the relationship between social phobia and avoidant personality disorder (see, for example, references 44–47). Our results suggest that from a genetic perspective, social phobia belongs with avoidant personality disorder on axis II.

Our results supporting a genetic distinction between axis I and axis II disorders might be seen as surprising, given previous evidence that they are highly comorbid and hard to distinguish empirically (48). Another plausible interpretation of our findings, particularly for internalizing disorders, is that different sets of genetic risk factors predispose to psychiatric disorders that are typically transient and episodic in nature and those that are characteristically more chronic.

Third, “transitional” disorders with substantial loadings on two genetic factors provide further insights into the structure of the genetic risk for psychiatric disorders. Furthermore, the existence of these transitional disorders indicates that the psychiatric disorders in our current classification do not neatly fall into our four proposed clusters. Individuals with high criterion counts for borderline personality

* Four factors were identified using an oblique geomin rotation. Only those paths are depicted that account for more than 10% of the genetic variance in disease liability—that is, have a path estimate of ≥0.316. For all path estimates in the model, see Table 3. PD=personality disorder.
axis I externalizing disorders. Environmentally, borderline personality disorder has links with all personality disorders and with axis I internalizing disorders; genetically, it is closely tied to axis I and II externalizing disorders. Consistent with our earlier study (13), the division of common psychiatric disorders into internalizing versus externalizing factors results from genetic and not from environmental risk factors. By contrast, the division into axis I versus axis II disorders arises from the effects of both genes and the environment.

Our results are also congruent with our previous analysis in this sample of the structure of genetic risk factors for personality disorders (27). That study, which examined only the 10 personality disorders, identified three genetic factors, the first of which loaded most heavily on histrionic, narcissistic, and borderline personality disorders—clearly reflecting our axis II externalizing factor. The second factor loaded more specifically on antisocial and borderline personality disorders—approximating our broader axis I externalizing genetic factor. The third genetic factor loaded most heavily on avoidant and schizotypal personality disorders, with weaker loadings on dependent and schizoid personality disorders—reflecting our axis II internalizing genetic factor.

**Limitations**

These results need to be interpreted in the context of nine additional potentially significant limitations. First, our
results are obtained in native-born young adult Norwegian twins and may not generalize to other ethnic or age groups.

Second, as many important psychiatric disorders (e.g., schizophrenia, autism, bipolar illness) were not included in these analyses, no claims can be made for our identification of the structure of risk factors for all psychiatric illness.

Third, using traditional statistical methods, we were unable to estimate results separately in male and female participants. While we controlled for prevalence differences across the sexes, we cannot rule out the possibility that we have averaged results of the two sexes that might meaningfully differ from one another. However, three findings reduce our concern that we have thereby introduced significant biases in our findings. In our earlier multivariate study in the Virginia Twin Registry (13), once we accounted for differences in prevalence, we were able, in a much larger twin sample, to constrain to equality parameter estimates across the sexes. In all of our previous analyses of the axis I and II disorders in this Norwegian sample, we have failed to find evidence for sex-specific genetic or environmental effects (17–19, 27, 49–52). Finally, we examined several models of our 22 disorders treating the criterion counts and subthreshold and threshold diagnoses as normally distributed variables. While this approach does not correctly capture the distributional properties of our variables, it nonetheless provides some useful information. Compared to the full model with separate parameter estimates for male and female participants, a model constraining all the genetic and environmental parameters to equality in the two sexes provided a much better fit using the Bayesian information criterion, a fit index particularly well suited for complex models (53).

Fourth, we were unable to test, using standard twin model fitting, whether the addition of shared environmental factors would improve the fit of this large multi-
variate model. However, a wide range of previous analyses with most of the disorders included in our model failed to find evidence for substantial shared environmental effects (17–19, 27, 47, 49–52, 54). Furthermore, treating the criterion counts and subthreshold and threshold diagnoses as normally distributed, we compared the full model and a model that dropped all of the shared environmental parameters. It fitted much better using the Bayesian information criterion and also was clearly superior to a model that dropped all the additive genetic parameters. While we cannot rule out a modest degree of confounding of genetic with shared environmental effects, it is unlikely that this confounding is substantial.

Fifth, we could not formally test the number of genetic and environmental factors extracted. We therefore had to rely on the more traditional methods of the scree plot and clinical interpretation. We feel confident, however, that four genetic and three specific-environmental factors represent the most parsimonious structure that well accounts for the observed results.

Sixth, we lacked the ability to calculate confidence intervals for the individual parameter estimates. Given the size of our sample, we suspect that our parameter estimates are known with only moderate accuracy (55). However, it is the broad pattern of our findings rather than the specific value of any individual parameter that is probably of greatest value in these analyses.

Seventh, substantial attrition was observed from the original birth registry through three waves of contact. However, detailed analyses of the predictors of nonresponse across waves (26) revealed that cooperation was strongly predicted by sex, zygotici, age, and education but not psychiatric symptoms or self-report personality disorder items that have been shown empirically to predict DSM-IV personality disorder criteria in the personal interview phase. For example, among 45 predictors, including 22 mental health variables, only two—older age and monozygosity—predicted cooperation in the personal interview phase. Twin analyses of 25 mental health-related variables from earlier questionnaires reflecting psychiatric and personality disorder symptoms and substance use revealed no significant differences between those who completed a personal interview and those who did not (26). Thus it is unlikely that attrition introduced bias in the estimates of the etiological role of genetic and environmental risk factors for this broad range of mental health indicators. Our sample is probably broadly representative of the Norwegian population with respect to psychopathology.

Eighth, could our results be sensitive to the specific method of factor extraction? In addition to the oblique geomin rotations, we examined solutions obtained by the orthogonal varimax and oblique promax methods. All four genetic factors and the first environmental factor were stable across rotational methods, with only small differences on the second and third environmental factors (e.g., higher cross-loadings for antisocial personality disorder and drug abuse or dependence). The main features of our results, especially the four genetic factors, were stable with respect to the method of factor extraction.

Finally, could method variance account for critical parts of our findings? We used two separate instruments with different formats for the assessment of axis I versus conduct and personality disorders. However, our results suggest that this concern is unwarranted. Our second genetic factor contained five axis II personality disorders and two axis I disorders. Antisocial personality disorder and conduct disorder, both assessed in our personality disorder interview, were placed in the third genetic factor and the third environmental factor each time with other axis I disorders. This pattern of results is not consistent with a method variance account.

Conclusions

We have long sought to ground our diagnostic categories in etiological processes. While not free of methodological limitations, this study nonetheless provides, for the first time, a comprehensive picture of the structure of genetic and environmental risk factors for most common DSM-IV axis I disorders and all axis II disorders. The genetic structure of these disorders, as indexed by syndromal and subsyndromal diagnoses of the axis I disorders and criterion counts for the personality disorders, was relatively simple, consisting of four factors reflecting the two major dimensions of internalizing versus externalizing and axis I versus axis II. However, for five disorders, substantial genetic influences from two of these factors were required to adequately explain their etiology. The structure of the environmental influences on these disorders, by contrast, looked quite different, indicating that the organization of common psychiatric disorders into these coherent groups was largely a result of genetic and not environmental influences. Our findings reinforce accumulating evidence over the past half century that genetic factors play a critical role not only in the etiology of individual disorders but also in the structure of disorders as they occur and co-occur in human populations.
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