Gambling disorder (GD), characterized by problematic gambling leading to significant impairment, has recently been re-classified in the Diagnostic and Statistical Manual of Mental Disorders, Version 5 (DSM-5) (APA, 2013) as belonging to the same category as the substance use disorders (SUDs). This was based on emerging evidence of shared neurobiological underpinnings and phenomenology. In contrast to SUDs, however, there has been little research examining contributions of genetic and environmental influences to risk for GD (Slutske et al., 2010; Slutske et al., 2013).

There have been two major twin studies of GD to date (Eisen et al., 1998; Slutske et al., 2010). The larger of the studies included 3359 all-male pairs from the national United States Vietnam-Era Twin Registry (VET; Eisen et al., 1998). The results of fitting biometric structural equation models to the VET data yielded estimates of the percentage of variation in DSM-III-R (APA, 1987) GD liability explained by genetic, shared environmental, and unique environmental factors of 46, 16, and 38%, respectively, although the source of environmental contributions or sex differences in GD liability.

In the replication study, when using a lower GD threshold, there was evidence for significant genetic (60%; 95% confidence interval (CI) 45–76%) and unique environmental (40%; 95% CI 24–56%), but not shared environmental contributions (0%; 95% CI 0–0%) to GD liability; this did not significantly differ from the original study. In the combined analysis, higher GD thresholds (such as one consistent with DSM-5 GD) and a multiple threshold definitions of GD yielded similar results. There was no evidence for quantitative or qualitative sex differences in the liability for GD.

Conclusions. Twin studies of GD are few in number but they tell a remarkably similar story: substantial genetic and unique environmental influences, with no evidence for shared environmental contributions or sex differences in GD liability.
and Australia), >10 years apart, and with different gender compositions, the results were strikingly similar. Both studies arrived at the conclusion that about half of the variation in GD liability was explained by genetic factors, that the shared environment did not contribute significantly to variation, and that unique environmental factors accounted for the remaining variation.

More recently, two smaller twin studies of GD have been conducted. One study recruited 912 twin and sibling pairs via a web-based survey (Blanco et al., 2012). A broad lifetime measure of GD was created based on the number of gambling episodes and GD symptomatology. The estimate of genetic influences was 83% with no evidence for shared environmental influences or sex differences (Blanco et al., 2012). This estimate is higher than those found in previous twin studies of GD (Eisen et al., 1998; Slutske et al., 2010) and may be due to the low prevalence rates of GD observed or the different operationalization of GD compared with previous studies.

The other study was based on data collected within the context of an ongoing longitudinal twin study in Minnesota (King et al., 2017). At ages 18 and 25, participants completed questionnaires of past-year symptoms of GD as measured by the South Oaks Gambling Screen (Lesieur and Blume, 1987). Estimates of the contribution of genetic factors to variation in GD were 5–37%, varying by age and sex, with genetic contributions increasing over time for men and decreasing over time for women. Additionally, there was evidence of significant shared environmental influences (19–29%) at age 18 (King et al., 2017). These discrepant findings might be due to the relative youth of this sample, the low yield of gambling problems, or the focus on past-year rather than lifetime pathology.

Collectively, these four twin studies of GD included just 7916 twin pairs. By comparison, meta-analytic reviews have identified rather than lifetime pathology. Discrepant findings might be due to the relative youth of this sample, the low yield of gambling problems, or the focus on past-year rather than lifetime pathology.

The current study represents an attempt to replicate the results of the ATR study (Slutske et al., 2010) in a new independent Australian twin cohort. In addition to attempting to replicate the previous results in a new sample, we conducted a more powerful analysis by combining data from the original ATR study with the new Australian twin cohort in an effort to achieve the requisite sample size to detect significant sex differences or shared environmental variation in liability for these addictive disorders (Verweij et al., 2010; Verhulst et al., 2015).

The current study was extensive, with many detailed questions about gambling involvement prior to the assessment of GD symptoms.

Participants

Replication study
Participants were 3292 members of the ATR Cohort III. The sample included adult twins born between 1972 and 1979. The data were collected by computer-assisted telephone interviews conducted between 2005 and 2009 (individual response rate of 76%). There were 1205 complete twin pairs (565 MZ [396 female, 169 male], 640 DZ [299 female–female, 116 male–male, and 225 female–male]), and 882 individual twins from incomplete pairs (321 MZ [180 female, 141 male], 561 DZ [136 female–female, 136 male–male, and 289 female–male]). The mean age was 31.8 years (range = 27–40) and 64% of the sample was female. Further details are reported in Lynskey et al. (2012).

Original study
Participants were 4764 members of the ATR Cohort II. The sample included adult twins born between 1964 and 1971. In 2004–2007, a telephone interview containing a thorough assessment of gambling behaviors was conducted with the ATR Cohort II members (individual response rate of 80%). The mean age was 37.7 years (range = 32–43) and 57.2% of the sample was female. There were 1875 complete twin pairs (867 MZ [520 female, 347 male], 1008 DZ [367 female–female, 227 male–male, and 414 female–male]), and 1014 individual twins from incomplete pairs (304 MZ [151 female, 153 male], 710 DZ [181 female–female, 216 male–male, and 313 male–female]). Further details about the study participants are reported in Slutske et al. (2010).

Procedure

Twins were assessed by structured telephone interviews. Interviews were administered by trained lay interviewers who were blind to the status of the cotwin. Interviewers were supervised by project editors, who reviewed all interview protocols.

Although both studies were based on national Australian samples, there were differences in the state or territory of residence of the participants ($\chi^2 = 119.82, df = 7, p < 0.0001$). This is important because the eight different states/territories differ in their densities of gambling venues (Productivity Commission, 2010), which has been linked to the prevalence of gambling problems (Productivity Commission, 1999). Compared with the original study, the replication study had a larger percentage of participants hailing from the state with the greatest densities of venues (New South Wales; 23.3% v. 19.6%) and a smaller percentage from the state with the lowest gambling venue density (Western Australia; 12.6% v. 14.7%).

Another potentially important difference between the two studies was the extent to which gambling was the focus of the recruitment and assessment. The primary focus of the replication was cannabis use, whereas the focus of the original study was gambling involvement. Therefore, recruitment materials and supporting documentation available to prospective participants described the replication study as a ‘cannabis’ study and the original study as a ‘gambling’ study. The gambling assessment in the ‘cannabis’ replication study was brief, including the 10 DSM-IV GD symptoms and two screening questions about the extent of gambling involvement; the gambling assessment in the original ‘gambling’ study was extensive, with many detailed questions about gambling involvement prior to the assessment of GD symptoms.

Notes
†The notes appear after the main text.
All interviews were tape-recorded and a random sample of 5% of the interview tapes was reviewed for quality control. The replication study was approved by the Institutional Review Boards at Washington University and Berghofer QIMR, and secondary analysis of the data was approved by the University of Missouri. The original study was approved by the Institutional Review Boards at the University of Missouri and QIMR. All participants provided informed consent.

**Measures**

The same GD assessment was used in the two studies: the NORC DSM-IV Screen for Gambling Problems (NODS; Gerstein et al., 1999). The DSM-IV diagnostic criteria for GD are primarily composed of symptoms modeled on the substance dependence criteria, including the concepts of preoccupation, loss of control, tolerance, and withdrawal, along with two symptoms related to legal and financial consequences of gambling. One symptom that is unique to GD is the concept of ‘chasing losses,’ that is, following gambling losses with more gambling to ‘get even.’ A diagnosis of DSM-IV GD requires endorsing five of 10 diagnostic criteria. The DSM-IV diagnostic criteria were assessed for all participants who reported they had ever gambled at least five times within a 12-month period. Those participants who did not endorse this item were considered to be asymptomatic with respect to GD diagnostic criteria. Although both studies pre-dated the release of the DSM-5, it was possible to derive a DSM-5 GD diagnosis from the data by eliminating the one symptom that was omitted from the DSM-5 criteria (committed illegal acts to finance gambling) and lowering the number of symptoms required from five to four. The test-retest reliability of DSM-5 GD in the original cohort was very good (kappa = 0.75; Yule’s Y = 0.82; Slutske et al., 2013).

**Analytic plan**

Survey analysis procedures in SAS (SAS Institute, 2015) were used to test whether there were differences in the prevalences of gambling involvement and disorder in the two twin cohorts while taking into account the non-independence of twin pair observations. Differences in the proportions of men and women in the two samples were accounted for by regressing out the effect of sex on the outcome of interest.

Twin correlations in GD liability were estimated in Mplus (Muthén and Muthén, 2017). Within the replication sample, evidence for genetic influences was tested by comparing the same-sex MZ and DZ twin correlations. As MZ twins share all of their DNA and DZ twins share just half of their segregating DNA, higher MZ twin correlations compared with DZ twin correlations would provide evidence for genetic influences on GD symptomatology. Quantitative sex differences, or differences in the magnitude of genetic and environmental influences, were tested by comparing the within-zygosity differences in the twin correlations obtained from the same-sex male v. female twin pairs. Finally, qualitative sex differences were tested by comparing the twin correlations in unlike-sex and same-sex dizygotic twin pairs. If there are different genetic factors influencing liability in men and women, or qualitative sex differences, it would be expected that the same sex DZ twin pairs would have higher twin correlations than unlike-sex pairs.

Biometric models were fit directly to the raw twin data using robust weighted least squares within Mplus (Muthén and Muthén, 2017). Liability threshold models, which assume a latent liability continuum underlying a categorical diagnosis, were fit to the twin data (Neale and Cardon, 1992; Kendler, 1993). Biometric model-fitting was conducted to partition the variation in GD liability into additive genetic (A), shared environmental (C), and unique environmental influences (E; estimates of unique environmental variation also include measurement error). Biometric model-fitting uses the known relationship between twins to partition resemblance into genetic factors (A), which contribute twice as much to resemblance in MZ compared with DZ twins, and shared environmental factors (C), which contribute equally to resemblance in MZ and DZ twins. In addition, the biometric model contains individual-specific factors (E), which reflect the impact of environmental experiences specific to one twin that may lead to differences in liability for GD. Analyses were first conducted in the replication sample based on a GD threshold of 1+ symptoms. When the replication and original samples were subsequently combined, four different thresholds were examined: 1+, 2+, 3+, or 4+ symptoms of GD (consistent with the DSM-5 GD diagnosis). A three-level multiple-threshold model was also fit, with thresholds of no symptomatology, subthreshold symptomatology (1–3 GD symptoms), and suprathreshold symptomatology consistent with a GD diagnosis (4+ GD symptoms).

Evidence for quantitative sex differences was evaluated by comparing the fit of a model in which estimates of the variance components were constrained to be equal in men and women to a model that allowed estimates for men and women to vary. A significant decrease in model fit would indicate the presence of quantitative sex differences. Evidence for qualitative sex differences was tested by comparing the fit of a model in which the genetic correlation for the unlike-sex twin pairs was set to 0.5 (the genetic correlation for the same-sex dizygotic twin pairs) to a model in which it was freely estimated. A significant decrease in model fit would indicate the presence of qualitative sex differences. Evidence for cohort differences was evaluated by comparing the fit of a model in which the estimates of the variance components were constrained to be equal in the replication and original twin cohorts to a model that allowed the estimates in the two twin cohorts to vary. Model comparisons were conducted using the Satorra–Bentler scaled χ² difference test (Satorra and Bentler, 2010).

**Results**

**Prevalences**

**Replication sample**

Fifty percent of participants had gambled at least five times in a single year and 11% had gambled at least once a week (see Table 1). The overall lifetime prevalence of GD based on DSM-5 criteria was 2.07% (4.08% among men, 0.98% among women; Table 1). The overall lifetime prevalence of ever experiencing one or more DSM-5 GD symptoms was 7.8% (14.3% among men, 4.3% among women). Prevalence rates of experiencing one or more GD symptoms were similar for those with an unlike-sex DZ co-twin compared with those with a same-sex DZ co-twin (men: unlike-sex = 9.0% v. same sex = 9.1%; women: unlike-sex = 3.1% v. same sex = 3.2%).

**Original sample**

The prevalences of gambling at least five times in a single year (t = −22.53, df = 4974, p < 0.0001), gambling at least once a week (t = −21.77, df = 4974, p < 0.0001), and experiencing one or more GD symptoms (t = 5.44, df = 4969, p < 0.0001) were
all significantly lower in the replication than in the original sample. The prevalences of DSM-5 GD in the two cohorts did not significantly differ ($t = -1.29, df = 4974, p = 0.20$). (See online Supplemental Materials for an examination of the cohort differences in gambling involvement prevalence.)

**Twin correlations**

**Replication sample**

Twin correlations in liability for 1+ GD symptoms provided some initial indication for genetic contributions to liability for GD. The correlations in GD liability were higher among the MZ twins, who share all the same genes, compared with the DZ twin pairs, who share just half of their segregating genes, in both men and women (Table 2); however, these correlations did not significantly differ ($\Delta \chi^2 = 2.64, df = 2, p = 0.27$). There was no evidence for quantitative sex differences as twin correlations for men and women within the same-sex zygosity groups did not differ significantly ($\Delta \chi^2 = 2.38, df = 2, p = 0.31$). The low unlike-sex twin correlation ($r = 0.05; 95\%$ confidence interval (CI) $-0.47$ to $0.56$) is of note, as this correlation suggests a potential qualitative sex difference. However, twin correlations for unlike-sex and the same-sex dizygotic twin pairs did not significantly differ ($\Delta \chi^2 = 2.64, df = 2, p = 0.27$).

**Combined sample**

Twin correlations for the original sample were similar to those obtained for the replication sample (Table 2), and correlations in the two cohorts did not significantly differ ($\Delta \chi^2 = 3.58, df = 5, p = 0.61$). When combining the original and replication cohorts, MZ twin correlations remained higher than DZ twin correlations for both men and women; with the increased power of the combined sample, these differences were now significant ($\Delta \chi^2 = 9.56, df = 2, p = 0.01$). No evidence was found for quantitative ($\Delta \chi^2 = 1.52, df = 2, p = 0.47$) or qualitative ($\Delta \chi^2 = 0.18, df = 2, p = 0.91$) sex differences using the combined sample.

**Biometric model fitting**

**Replication sample**

Biometric models were fit examining contributions to liability for experiencing 1+ GD symptoms (Table 3). In these and all subsequent biometric models, thresholds for men and women were allowed to vary because they could not be constrained to be equal without a significant deterioration in model fit ($\Delta \chi^2 = 83.52, df = 1, p < 0.0001$). The best fitting model was one that included additive genetic and unique environmental sources of variation – shared environmental factors did not account for a significant portion of the variation in liability for GD. There was no evidence of quantitative sex differences, as parameter estimates did not differ significantly for men and women ($\Delta \chi^2 = 5.21, df = 2, p = 0.07$). When constraining estimates to be equal for men and women, the heritability of GD liability was 60.3%, with unique environmental influences accounting for the remainder of the variance. There was no evidence for qualitative sex differences in that an unlike-sex twin genetic correlation of 0.50 could not be ruled out ($\Delta \chi^2 = 1.32, df = 1, p = 0.25$).

**Combined sample**

The results of the biometric model examining contributions to 1+ GD symptoms in the replication sample did not significantly differ from those obtained in the original sample; constraining parameter estimates across cohorts did not result in a significant decrease in model fit ($\Delta \chi^2 = 2.67, df = 4, p = 0.61$). The estimates of genetic, shared, and unique environmental influences in the combined sample were 45.6, 1.6, and 52.8%, respectively, among men, and 58.2, 0, and 41.7%, respectively, among women. However, these parameter estimates did not significantly differ ($\Delta \chi^2 = 1.23, df = 2, p = 0.54$). There was also no evidence for qualitative sex differences ($\Delta \chi^2 = 0.45, df = 1, p = 0.50$). The combined sample results after constraining the parameter estimates in men and women are presented in Table 4.

With the increased power from the combined sample, analyses were conducted examining influences on higher levels of GD symptomatology, including a threshold of 4+ GD symptoms, which is consistent with the DSM-5 GD diagnosis, and a three-level variable within a multiple-threshold model (Tables 2 and 4). Estimates obtained using the narrower 4+ symptoms’ threshold were similar to those obtained for the threshold of 1+ symptoms of GD, but with extremely broad CIs. Results of fitting the multiple-threshold model also yielded similar findings with narrower CIs. There was still no evidence

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**Table 1.** Lifetime prevalence of gambling involvement and GD among 8056 adult Australian twins from two independent cohorts

<table>
<thead>
<tr>
<th>Frequency of gambling</th>
<th>Cohort II (original study)</th>
<th>Cohort III (replication study)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Men $N = 3037$</td>
<td>Women $N = 2727$</td>
</tr>
<tr>
<td></td>
<td>% ($N$)</td>
<td>% ($N$)</td>
</tr>
<tr>
<td>Ever gambled 5+/year</td>
<td>81.3 (1655)</td>
<td>74.5 (2031)</td>
</tr>
<tr>
<td>Ever gambled weekly$^a$</td>
<td>39.1 (796)</td>
<td>33.7 (928)</td>
</tr>
<tr>
<td><strong>GD</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4+ GD symptoms$^b$</td>
<td>4.2 (86)</td>
<td>1.7 (45)</td>
</tr>
<tr>
<td>3+ GD symptoms</td>
<td>5.8 (118)</td>
<td>2.4 (64)</td>
</tr>
<tr>
<td>2+ GD symptoms</td>
<td>8.7 (178)</td>
<td>3.5 (95)</td>
</tr>
<tr>
<td>1+ GD symptoms</td>
<td>18.2 (370)</td>
<td>8.3 (225)</td>
</tr>
</tbody>
</table>

DSM-5, Diagnostic and Statistical Manual of Mental Disorders, Version 5.

$^a$Ever gambled at least once a week for at least six months in a row.

$^b$Corresponds to a DSM-5 diagnosis of GD.
Table 2. Twin correlations in liability for GD

<table>
<thead>
<tr>
<th>Threshold</th>
<th>MZ male</th>
<th>DZ male–male</th>
<th>MZ female</th>
<th>DZ female–female</th>
<th>DZ male–female</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Cohort II (original study)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1+ GD symptoms</td>
<td>0.49 (0.31–0.67)</td>
<td>0.26 (0.00–0.52)</td>
<td>0.52 (0.31–0.72)</td>
<td>0.26 (–0.05 to 0.58)</td>
<td>0.23 (0.02–0.45)</td>
</tr>
<tr>
<td><strong>Cohort III (replication study)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1+ GD symptoms</td>
<td>0.40 (0.12–0.69)</td>
<td>0.20 (0.06–0.35)</td>
<td>0.69 (0.50–0.88)</td>
<td>0.35 (0.25–0.44)</td>
<td>0.05 (–0.47 to 0.56)</td>
</tr>
<tr>
<td><strong>Cohorts II and III combined</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4+ GD symptoms*</td>
<td>0.59 (0.28–0.89)</td>
<td>0.32 (–0.08 to 0.72)</td>
<td>0.60 (0.32–0.88)</td>
<td>0.35 (–0.29 to 1.00)</td>
<td>0.03 (–1.00 to 1.00)</td>
</tr>
<tr>
<td>3+ GD symptoms</td>
<td>0.61 (0.38–0.84)</td>
<td>0.25 (–0.10 to 0.60)</td>
<td>0.71 (0.52–0.91)</td>
<td>0.21 (–0.67 to 1.00)</td>
<td>–0.03 (–1.00 to 1.00)</td>
</tr>
<tr>
<td>2+ GD symptoms</td>
<td>0.56 (0.37–0.76)</td>
<td>0.50 (0.27–0.74)</td>
<td>0.66 (0.48–0.85)</td>
<td>0.02 (–0.98 to 1.00)</td>
<td>–0.07 (–0.47 to 0.32)</td>
</tr>
<tr>
<td>1+ GD symptoms</td>
<td>0.47 (0.32–0.63)</td>
<td>0.25 (0.03–0.47)</td>
<td>0.60 (0.45–0.74)</td>
<td>0.18 (–0.10 to 0.46)</td>
<td>0.19 (0–0.38)</td>
</tr>
</tbody>
</table>

Note: Cell entries are tetrachoric correlations; 95% CIs are in parentheses. Bold typeface indicates a significant correlation.

*Corresponds to a DSM-5 diagnosis of GD.

Table 3. Parameter estimates of additive genetic (A), shared environmental (C), and unique environmental (E) influences from univariate biometric model-fitting of liability for GD in Cohort III (replication study)

<table>
<thead>
<tr>
<th>Parameters</th>
<th>A (95% CI)</th>
<th>C (95% CI)</th>
<th>E (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Full samplea</td>
<td>60.4 (44.5–76.3)</td>
<td>0.00 (0.0–0.0)</td>
<td>39.6 (23.7–55.5)</td>
</tr>
<tr>
<td>Men</td>
<td>40.3 (11.7–64.3)</td>
<td>0.00 (0.0–0.2)</td>
<td>59.6 (31.0–83.6)</td>
</tr>
<tr>
<td>Women</td>
<td>68.9 (49.8–84.9)</td>
<td>0.00 (0.0–0.1)</td>
<td>31.0 (11.9–47.0)</td>
</tr>
</tbody>
</table>

Note: 95% CIs are in parentheses.

*aUsing a threshold of 1+ GD symptoms.

for quantitative (Δχ² = 1.86, df = 2, p = 0.39) or qualitative sex differences (Δχ² = 0.53, df = 1, p = 0.47). The estimate of the genetic correlation between men and women was well below 0.50 for every GD threshold examined, but CIs were very broad and included 0.50 (Table 4).

Discussion

The findings of a previous Australian twin study (Slutske et al., 2010) were replicated in an independent Australian twin cohort. There was evidence for significant genetic (60%) and unique environmental (40%) contributions to GD liability, and despite the different levels of gambling involvement in the two cohorts, the relative contributions of genetic and environmental factors to GD liability did not significantly differ. In both the original and the replication samples, there was no evidence for a significant effect of the shared environment or for quantitative or qualitative sex differences in GD liability. Because twin studies of categorical phenotypes such as a GD diagnosis are often underpowered to detect such effects, we revisited their evidence after combining the two twin cohorts. In the combined sample – the largest twin study of GD to date – there was still no evidence for shared environmental influences or for quantitative or qualitative differences in the contribution of genetic and environmental factors to GD liability in men and women. These results do not appear to be a peculiarity of Australia, because lack of evidence for shared environment (Eisen et al., 1998; Blanco et al., 2012) and quantitative and qualitative sex differences (Blanco et al., 2012) have also been observed in national United States samples.

Evidence for sex differences

Prevalence rates of GD were significantly lower among women than among men, which is consistent with prior research (Petry et al., 2005). Additionally, although estimates of the heritability of GD in men and women in the combined sample did not significantly differ, they were somewhat disparate (46% among men v. 58% among women), suggesting a potential quantitative sex difference. Similarly, although the genetic correlation in GD liability of 0.35 in unlike-sex twin pairs was not significantly different from 0.50, it was suggestive of a potential qualitative sex difference. There are several lines of evidence suggesting it might be premature to rule out the possibility of differences between men and women in the genetics of GD. Men and women differ in the types of gambling they typically engage in, with men more likely to participate in strategic forms and women more likely to participate in non-strategic forms of gambling (Odlaug et al., 2011; Savage et al., 2014). These different forms of gambling appear to have distinct personality (Savage et al., 2014) and neurobiological correlates (van Holst et al., 2010). Men and women also differ in their performance on a gambling task [that simulates patterns of disadvantageous decision-making characteristic of GD (Brevers et al., 2013)], which is also related to neurobiological differences (van den Bos et al., 2013). In sum, there may be distinct etiologies of GD in men and women that were not detectable in this study.
Evidence for shared environmental contributions

In contrast, evidence ruling out an important influence of the shared environment was more conclusive in that the estimate was zero in the replication as well as the combined sample. Lack of evidence for a significant contribution of the shared environment may be due to the existence of gene–environment correlation (rGE) and gene–environment interaction (GxE). Depending on whether they involve an aspect of the shared or unique environment, these would be included in the estimates of ‘A’ or ‘E’, respectively. A study based on data from the original ATR Cohort II provided evidence for both rGE and GxE (Slutske et al., 2013). The genetic variation associated with the frequency of gambling was associated with exposure to neighborhood disadvantage (rGE), and the genetic risk associated with GD was associated with greater sensitivity to the deleterious effects of living in a disadvantaged neighborhood (GxE). There are likely many other genetically contingent environmental effects for GD yet to be discovered that should be explored in future research.

Evidence for genetic contributions

Given consistent evidence from twin studies demonstrating an important aggregate influence of genetic factors in the risk for GD among both men and women, a major challenge ahead will be to identify specific genetic variants that confer this risk. Unfortunately, molecular genetic research on GD is lagging even further behind than the quantitative genetic (twin) research. There have been only two genome-wide association studies (GWAS) of GD including a total of only 2742 participants, with no genome-wide significant single-nucleotide polymorphisms (SNPs) or genes detected in either study (Lind et al., 2013; Lang et al., 2016). The best clues so far come from (1) a series of reports on the incidence of GD among individuals with Parkinson’s disease (e.g. Weintraub et al., 2010) and restless legs syndrome (e.g. Tippmann-Peikert et al., 2007) being treated with a dopamine agonist medication that typically demonstrates relative selectivity for dopamine D3 receptors (Dodd et al., 2005; Tippmann-Peikert et al., 2007), (2) the replication in a rat model of an association of the dopamine D3 receptor gene with GD in humans (Lobo et al., 2015), and (3) a genome-wide significant association between risk-taking (a transdiagnostic endophenotype for GD) and a SNP in the CADM2 gene on chromosome 3 in a sample of over 116,000 individuals from the UK Biobank cohort (Strawbridge et al., 2018). As with other psychiatric disorders, progress in revealing the genetic underpinnings of GD will require worldwide cooperation and collaboration to amass the sample sizes required to detect genes of very small effect (Psychiatric GWAS Consortium Steering Committee, 2009; Sullivan et al., 2018). Because GWAS results can be extremely useful for more than identifying individual variants (Chabris et al., 2015; Maier et al., 2018), a top priority for the future will be to conduct more GWAS of GD.

Limitations

The main limitation of this study is that participants in the two twin cohorts represented a narrow range of ages (27–43 years), were primarily of Northern European ancestry, and resided in Australia – the results may not generalize to other ages, ethnicities or racial groups, or countries. This limitation is counterbalanced by the advantage of conducting this research in Australia. One of the greatest challenges to conducting community based twin studies of GD is the fact that it is relatively rare. In contrast to other countries such as the United States, most individuals in Australia have been heavily exposed to gambling opportunities, and Australia has one of the highest prevalences of GD in the world (Slutske et al., 2009). There are few other settings where this research could have been conducted.

Conclusions and implications

Consistent with previous research (Eisen et al., 1998; Slutske et al., 2010; Blanco et al., 2012), the current study suggests that genetic influences may be more important than shared environmental influences in the development of GD. An implication of these findings is that any explanation of the intergenerational transmission of gambling behavior that assumes that the intergenerational transmission is due to family environment rather than genetic transmission will be incomplete (Dowling et al., 2017). Efforts to prevent the intergenerational transmission of problematic gambling by targeting parental gambling may be misguided.

Replicable findings such as these increase confidence in scientific knowledge and lead to useful applications in society (Simons, 2014), including prevention and treatment efforts. For example, with increasing knowledge of the genetic and environmental underpinnings of tobacco use, there is interest in the potential

Table 4. Parameter estimates from combined analyses of Cohorts II and III (original and replication studies) of the contribution of additive genetic (A), shared environmental (C), and unique environmental (E) influences to the liability for GD at different thresholds

<table>
<thead>
<tr>
<th>Threshold</th>
<th>Parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>A (95% CI)</td>
</tr>
<tr>
<td>Multiple thresholdsa</td>
<td>53.9 (44.6–63.2)</td>
</tr>
<tr>
<td>4+ GD symptomsb</td>
<td>57.7 (0.0–1.00)</td>
</tr>
<tr>
<td>3+ GD symptoms</td>
<td>68.0 (54.0–81.9)</td>
</tr>
<tr>
<td>2+ GD symptoms</td>
<td>65.1 (52.4–77.0)</td>
</tr>
<tr>
<td>1+ GD symptoms</td>
<td>53.1 (43.1–63.1)</td>
</tr>
</tbody>
</table>

Note: 95% CIs are in parentheses.
GD, gambling disorder; rA, correlation of genetic influences among unlike-sex dizygotic pairs.
aModel included thresholds at 1+ symptoms and 4+ symptoms.
bCorresponds to a DSM-5 diagnosis of GD. Estimates are from models in which A, C, and E parameters were constrained across sex, thresholds were allowed to vary for men and women, and the effect of birth cohort differences in thresholds were covaried.
to use genetic and environmental information to inform clinical treatment efforts to reduce or quit smoking successfully (Piasecki, 2006), and to tailor interventions to the genotypes of the patient (precision medicine; Chen et al., 2018). Identifying specific genes relevant to GD, as well as how these genes interact with environmental influences, will be critical for improving treatment and prevention of GD.

Notes
1 The sample sizes provided were sometimes the number of twin pairs, sometimes the number of individuals.
2 In a classic twin study it is not possible to estimate both shared environmental and non-additive genetic influences within the same model. In the replication sample, a model that included additive genetic, shared environmental, and unique environmental sources of variation ($\chi^2 = 5.99$, df = 8, p = 0.65) fit slightly better than one that included dominant genetic variation ($\chi^2 = 7.59$, df = 8, p = 0.47). Therefore, ACE rather than ADE models were fit throughout.
3 A previous paper (Beaver et al., 2010) claimed to find evidence for quantitative sex differences for GD, but closer inspection revealed that the phenotype was not actually a measure of GD, and the evidence for sex differences was based on a handful of extreme outliers and improper data analysis (Slutske and Richmond-Rakerd, 2014). A re-analysis of the data yielded no evidence for quantitative sex differences for gambling involvement (Slutske and Richmond-Rakerd, 2014).

Supplementary material. The supplementary material for this article can be found at https://doi.org/10.1017/S0033291718002325

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Ethical standards. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008.

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