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# Shared Genetic Risk between Eating Disorder- and Substance-Use-Related Phenotypes: Evidence from Genome-Wide Association Studies — Source link

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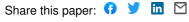
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# Shared genetic risk between eating disorder- and substanceuse-related phenotypes: Evidence from genome-wide association studies

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#### Abstract

Eating disorders and substance use disorders frequently co-occur. Twin studies reveal shared genetic variance between liabilities to eating disorders and substance use, with the strongest associations between symptoms of bulimia nervosa and problem alcohol use (genetic correlation  $[r_g]$ , twin-based = 0.23-0.53). We estimated the genetic correlation between eating disorder and substance use and disorder phenotypes using data from genome-wide association studies (GWAS). Four eating disorder phenotypes (anorexia nervosa [AN], AN with binge eating, AN without binge eating, and a bulimia nervosa factor score), and eight substance-use-related phenotypes (drinks per week, alcohol use disorder [AUD], smoking initiation, current smoking, cigarettes per day, nicotine dependence, cannabis initiation, and cannabis use disorder) from eight studies were included. Significant genetic correlations were adjusted for variants associated with major depressive disorder and schizophrenia. Total study sample sizes per phenotype ranged from ~2400 to ~537 000 individuals. We used linkage disequilibrium score regression to calculate single nucleotide polymorphismbased genetic correlations between eating disorder- and substance-use-related phenotypes. Significant positive genetic associations emerged between AUD and AN (r<sub>g</sub> = 0.18; false discovery rate q = 0.0006), cannabis initiation and AN ( $r_g$  = 0.23; q < 0.0001), and cannabis initiation and AN with binge eating ( $r_g$  = 0.27; q = 0.0016). Conversely, significant negative genetic correlations were observed between three nondiagnostic smoking phenotypes (smoking initiation, current smoking, and cigarettes per day) and AN without binge eating ( $r_{gs} = -0.19$  to -0.23; qs < 0.04). The genetic correlation between AUD and AN was no longer significant after co-varying for major depressive disorder loci. The patterns of association between eating disorder- and substance-use-related phenotypes highlights the potentially complex and substancespecific relationships among these behaviors.

#### **KEYWORDS**

eating disorders, genetic correlation, substance use

## 1 | INTRODUCTION

Well-established phenotypic associations exist between eating disorder and substance use phenotypes, with evidence for specific relations between particular types of eating disorders and substance use disorders. The prevalence of an alcohol use disorder (AUD) is greater among individuals with bulimia nervosa and binge-eating disorder than individuals with anorexia nervosa (AN) or healthy controls. <sup>1,2</sup> Similarly, individuals with bulimia nervosa or binge-eating disorder are at increased risk for smoking, nicotine dependence, <sup>3,4</sup> and cannabis use, <sup>4,5</sup> compared with individuals with AN or healthy controls, though

these results are not consistent.<sup>1</sup> Importantly, women with the binge-eating/purging subtype of AN report a higher prevalence of AUD, smoking, nicotine dependence, and cannabis use than women with the restricting subtype of AN.<sup>1,5,6</sup> Thus, binge eating—a transdiagnostic symptom defined as eating a large amount of food in a short period of time while experiencing loss of control—may be a key component of the observed association.

However, prior research has only partially addressed whether binge eating is the critical eating disorder symptom in the comorbidity, especially across different milestones of substance use (ie, initiation through substance use disorder) and across a variety of substances (ie, alcohol, nicotine, and cannabis). Elucidating shared sources for these associations is crucial because of the increased morbidity and mortality associated with comorbid presentations<sup>7,8</sup> and because improvements in one disorder may exacerbate (or weaken) symptoms of the other disorder.<sup>9</sup> Refining our understanding of these associations could improve prevention and treatment approaches for these debilitating disorders, their comorbidity, and their sequelae.

Accumulating findings from twin studies implicate shared genetic factors between eating disorder- and substance-use-related phenotypes. The strongest reported association is between bulimia nervosa symptoms (including binge eating) and problem alcohol use, with a genetic correlation  $(r_g)$  ranging from 0.23 to 0.53.<sup>10</sup> Although there has been less focus on genetic associations between bulimia nervosa symptoms and regular smoking and bulimia nervosa symptoms and illicit drug use disorder, twin-based  $r_{e}$ s of 0.35 and approximately 0.38, respectively, have been reported. 11,12 Limited information exists regarding whether less problematic aspects of substance use exhibit a significant  $r_g$  with eating disorder phenotypes. The impact of genetic factors influencing this comorbidity may significantly increase once an individual has progressed to problematic alcohol use, as genetic effects are more prominent in problem substance use, such as abuse and dependence, than with the initiation and general use of substances. 13-16 No study has comprehensively examined a range of eating disorder- and substance-use-related phenotypes to determine whether the  $r_g$  varies with different aspects of substance use and whether the  $r_g$  varies depending on the eating disorder and substance examined.

Recent advances in genomic methods allow for an assessment of  $r_g$  using existing genome-wide association study (GWAS) summary statistics. Unlike twin studies, these genome-wide methods allow for use of unrelated cases and controls, typically yielding sample sizes in the tens to hundreds of thousands. One such method, linkage disequilibrium score regression (LDSC),  $^{17,18}$  estimates single nucleotide polymorphism (SNP)-based heritability and  $r_g$  between phenotypes. Of particular relevance to low prevalence phenotypes, such as AN, estimation of SNP-based  $r_g$  does not require both phenotypes to be measured in the same individual; thus, independent studies assessing only one phenotype can be jointly examined.

The current study estimated SNP-based genetic correlations ( $r_g$ s) between eating disorder- and substance-use-related phenotypes based upon summary statistics from the largest published eating disorder GWAS and existing GWAS encompassing a range of substance-use-related phenotypes (ie, alcohol, nicotine, and cannabis), using robust data from twin studies to shape our three hypotheses. First, we hypothesized that the strongest SNP-based  $r_g$  would be between eating disorder phenotypes that have binge eating as a core symptom and alcohol use phenotypes, rather than between eating disorder phenotypes and nicotine and cannabis use-related phenotypes. <sup>10</sup> Second, we hypothesized that for binge eating-related phenotypes, the SNP-based  $r_g$  would be higher when assessing AUD than typical alcohol consumption, <sup>10</sup> because we expected that two problem behaviors are more likely to share genetic risk than a problem behavior (eg, binge eating) and a normative pattern (eg, alcohol consumption). Because

we have less information from twin studies about genetic associations between liabilities to eating disorders and tobacco (nicotine) and cannabis use-related phenotypes, we do not forward specific hypotheses for these substances. Finally, prior studies document robust genetic associations for major depressive disorder and schizophrenia with both eating disorders and substance-use-related phenotypes.  $^{19-21}$  We hypothesized that  $r_g$ s between eating disorders and substance use and disorder would be attenuated when accounting for variants associated with major depressive disorder and schizophrenia. Findings from this study will yield important information about the role of genetics in this clinically challenging pattern of comorbidity.  $^{22}$ 

#### 2 | METHOD

#### 2.1 | Participants

We included summary statistics from two existing GWAS of eating disorder phenotypes where participants were primarily of European ancestry<sup>21,23</sup> and data from individuals of European ancestry from six existing GWAS of substance-use-related phenotypes. 19,20,24-27 The eating disorder phenotypes (Table 1) included a diagnosis of AN (which was further parsed into AN with binge eating or AN without binge eating) and a bulimia nervosa factor score derived from the Eating Disorder Examination,<sup>28</sup> a well-established structured clinical interview for eating disorders. We did not examine bulimia nervosa or binge-eating disorder because there are currently no published GWAS for either disorder; thus, the bulimia nervosa factor score represents the closest to a GWAS of bulimia nervosa available. Substance-userelated phenotypes ranged from typical use (eg, drinks per week, smoking initiation, and cannabis initiation) to substance use disorder (ie, AUD, nicotine dependence, and cannabis use disorder). Sample sizes for the phenotypes ranged from 2442 (bulimia nervosa factor score) to 537 349 (drinks per week) individuals. Table 2 provides individual study details.

# 2.2 | Statistical analysis

We used LDSC<sup>17,18</sup> to evaluate SNP-based genetic correlations ( $r_g$ ) between samples. This method uses the linkage disequilibrium (LD) structure of the genome to estimate the distribution of effect sizes for individual SNPs as a function of their LD score. Under a polygenic model, causal SNPs are likely to be overrepresented in higher LD score bins (ie, including additional SNPs in high LD), such that associations with SNPs in these LD bins will make stronger contributions to the phenotypic variation under study. This polygenic distribution of effect sizes across LD score bins provides an estimate of SNP-based heritability, that is, the proportion of phenotypic variance that is attributable to the aggregate effects of genome-wide SNPs. The correlation of effect sizes across LD bins between two phenotypes then provides an estimate of SNP-based  $r_g$ .

Genetic correlations range from -1 to +1, where the sign indicates that the same genetic factors are contributing to variation in the target traits in *opposite* or *same* directions, respectively. The LDSC intercept for the genetic covariance provides evidence about sample overlap across two traits. SNPs (MAF > 0.01) found in the HapMap3 EUR population were used to calculate LD scores. We used the false discovery rate<sup>29</sup> to correct for multiple testing (n = 66 tests; q < 0.05). Finally, post hoc analyses examined whether significant differences between two  $r_g$ s existed, using the jackknife procedure implemented through LDSC.<sup>17</sup>

We used GNOVA $^{30}$  to stratify significant  $r_g s$  between the eating disorder- and substance-use-related phenotypes into both tissue-specific (for seven broadly defined tissue classes: brain, cardiovascular, epithelial, gastrointestinal, immune-related, muscular, and "other") and nontissue-specific functional regions of the genome. GenoCanyon $^{31}$  and GenoSkyline $^{32,33}$  annotation methods, which integrate transcriptomic and epigenomic data from ENCODE $^{34}$  and the Roadmap Epigenomics Project, $^{35}$  were used to define functional regions of the genome.

Finally, for significant  $r_g$ s detected in LDSC, multitrait-based conditional and joint analysis using GWAS summary data (mtCOJO)<sup>36</sup> was used to condition both input GWAS (eg, AN and AUD) for variants associated with major depressive disorder<sup>37</sup> at  $P < 5 \times 10^{-7}$  and schizophrenia<sup>38</sup> at  $P < 5 \times 10^{-8}$ . Because fewer genome-wide significant SNPs were identified for major depressive disorder than schizophrenia, we chose a more lenient P value threshold for major depressive disorder to capture a comparable number of SNPs. LDSC was used to compute  $r_g$ s using the resulting genome-wide summary statistics for each trait after separately adjusting for major depressive disorder or schizophrenia variants to examine whether conditioning on either disorder would affect the observed genetic relationships.

## 3 | RESULTS

The overall SNP-based heritability for the eating disorder phenotypes ranged from 0.20 to 0.39, whereas the corresponding heritabilities for the substance-use-related phenotypes ranged from 0.03 to 0.35 (Table S1). Figure 1 and Table S1 show the genetic correlations (r<sub>o</sub>s) between all four eating disorder phenotypes and eight substance-userelated phenotypes. Broadly speaking, there were significant  $r_{\sigma}$ s across substance-use-related phenotypes, ranging from 0.21 (AUD and cigarettes per day) to 0.70 (drinks per week and AUD). Cannabis initiation risk was not significantly genetically correlated with cigarettes per day or nicotine dependence. For the remaining results, we focus on previously unexplored associations of interest in this studycorrelations between eating disorder- and substance-use-related phenotypes. For these associations, the genetic covariance intercepts ranged from -0.03 (standard error [SE] = 0.01; AN and cannabis initiation) to 0.01 (SE = 0.01; AN and cannabis use disorder), indicating some sample overlap (or low-level confounding) existed, 39 although the LDSC approach parses this overlap from the  $r_g$  estimation.

**TABLE 1** Eating disorder-related phenotype descriptions

Phenotype	Definitions		
Anorexia nervosa (AN) <sup>a</sup>	Diagnostic criteria included the following:		
	1. Body mass index less than minimally expected		
	2. Intense fear of gaining weight		
	<ol><li>Weight or shape disturbance, undue influence of weight or shape, or denial of the seriousness of the disorder</li></ol>		
AN with binge eating <sup>b</sup>	Individuals with AN who also engaged in binge-eating episodes, defined as eating a large amount of food in a short period of time while having a sense of loss of control over the eating episode.		
AN without binge eating <sup>b</sup>	Individuals with AN who did not engage in binge-eating episodes.		
Bulimia nervosa (BN) <sup>c</sup> factor	Derived from a factor analysis that included the following items:		
	Reporting self-induced vomiting to control body weight		
	Reporting suffering from or being treated for binge eating		
	3. Reporting suffering from or being treated for bulimia		

<sup>a</sup>A fourth diagnostic criterion for AN includes amenorrhea. However, amenorrhea was excluded as a required criterion for cases in the Psychiatric Genomics Consortium datasets because it is no longer a diagnostic criterion in the DSM-5.

bThe DSM and ICD include two subtypes of anorexia nervosa (AN)—a binge-eating/purging subtype and a restricting subtype. Although it would have been ideal to examine differences between the AN binge-eating/purging subtype and AN restricting subtype, this was not possible with current Psychiatric Genomics Consortium data. However, there was sufficient information about presence or absence of binge eating, which resulted in creating the AN with binge-eating and AN without binge-eating subtypes.

<sup>c</sup>Bulimia nervosa is defined as (a) recurrent episodes of binge eating, (b) recurrent inappropriate compensatory behaviors (eg, self-induced vomiting or laxative use) to prevent weight gain, (c) the binge eating and inappropriate compensatory behaviors occurring an average of twice a week for 3 mo, (d) having undue influence of body weight and shape, and (e) disturbance not occurring during AN.

Significant positive  $r_g$ s were observed for alcohol- and cannabis use-related phenotypes. First, the  $r_g$  was significant between AN and AUD ( $r_g$  = 0.18; SE = 0.05; q = 0.0006) but not between AN and drinks per week ( $r_g$  = 0.01; SE = 0.03; q = 0.91), suggesting that genetic factors that increase risk for AN also increase risk for AUD, but little evidence exists for shared genetic risk between AN and typical alcohol consumption. These two correlations significantly differed from each other (z-score = 3.51, P = 0.0005). Intriguingly, there was a significant difference in  $r_g$ s for AN and AUD versus AN without binge eating and AUD (z-score = 2.28, P = 0.02) but not for AN and AUD versus AN with binge eating and AUD (z-score = 0.23, P = 0.82). The genetic covariance estimates between AN and AUD were significant in both functional (corrected  $\rho_g$  = 0.01; corrected r = 0.23; corrected r = 0.007) and nonfunctional categories (corrected  $\rho_g$  = 0.01; corrected r = 0.19; corrected q = 0.002; Table S2) but not in any specific tissue

type. No significant association between the bulimia nervosa factor score, which included items pertaining to both binge eating and compensatory behaviors, and either alcohol-use-related phenotype was observed.

Second, the significant  $r_g$  between AN and cannabis initiation was 0.23 (SE = 0.04, q < 0.0001) and the significant  $r_g$  between AN with binge eating and cannabis initiation was 0.27 (SE = 0.08, q = 0.0017), indicating that genetic factors that increase the risk for AN may also increase risk for cannabis initiation. However, cannabis initiation was not significantly correlated with the bulimia nervosa factor score ( $r_g$  = 0.15, SE = 0.18, q = 0.57) or with AN without binge eating ( $r_g$  = 0.10, SE = 0.08, q = 0.31). No significant associations were observed between any eating disorder phenotype and cannabis use disorder ( $r_g$ s

= -0.08-0.23; SEs = 0.01;  $qs \le 0.57$ ). Post hoc analyses revealed significant differences in the  $r_g$ s for AN and cannabis initiation versus AN and cannabis use disorder (z-score = 2.70, P = 0.01). However, the  $r_g$  between AN with binge eating and cannabis initiation, while significant, was statistically different from the  $r_g$  between AN with binge eating and cannabis use disorder. The genetic covariance estimate between AN with binge eating and cannabis initiation was significant in both functional (corrected  $\rho_g$  = 0.01; corrected r = 0.60; corrected q < 0.0001) and nonfunctional categories (corrected  $\rho_g$  = 0.01; corrected r = 0.30; corrected q = 0.004; Table S3) but not in any specific tissue type. The genetic covariance estimate between AN without binge eating and cannabis initiation was only significant in

**TABLE 2** Details of samples included in analyses

ADLE Z	Details of Samples inch	aucu iii aiiaiyses				
Study	Sample/Consortium	Phenotype(s)	Definition	Sample Size (cases/controls if binary)	Number of SNPs in Summary Statistics File	
Eating disorder phenotype						
Watson et al (2019)	PGC-ED	1. Anorexia nervosa	DSM-III-R, DSM-IV, ICD-8, ICD-9, ICD-10, or self-reported anorexia nervosa	16 992/55 525	8 219 102	
		2. Anorexia nervosa with binge eating		2381/10 249	8 982 440	
		3. Anorexia nervosa without binge eating		2262/10 254	8 671 192	
Wade et al (2013)	Australian Twin Registry	Bulimia nervosa factor	Eating Disorder Examination	151/2291	6 150 213	
Substance use-related phenotype						
Kranzler et al (2019)	MVP	Alcohol use disorder	ICD-9 or ICD-10	34 658/167 346	6 895 251	
Walters et al (2018)	PGC-SUD	Alcohol dependence	DSM-IV	8485/20 272	9 271 145	
Liu et al (2019)	GSCAN	1. Drinks per week <sup>a</sup>	Average number of drinks each week	537 349	11 916 707	
		2. Smoking initiation	Ever vs never regular smoker	311 629/321 173	11 733 344	
		3. Current smoking <sup>c</sup>	Current vs former smokers	92 573/220 248	12 197 133	
		4. Cigarettes per day <sup>a</sup>	Average number of cigarettes smoked per day	263 954	12 003 613	
Hancock et al (2017)	14 samples	Nicotine dependence <sup>b</sup>	Mild (FTND score 0-3)Moderate (FTND score 4-6)Sever (FTND score 7-10)	14 184 (Mild)	10 622 668	
				9206 (Moderate)		
				5287 (Severe)		
Pasman et al (2018)	ICC UK Biobank	Cannabis initiation	Lifetime cannabis use	43 380/118 702	11 733 371	
Demontis et al (2019)	iPSYCH	Cannabis use disorder	ICD-10	2387/48 985	8 969 939	

Abbreviations: DSM, Diagnostic and Statistical Manual; FTND, Fagerström Test of Nicotine Dependence; GSCAN, GWAS & Sequencing Consortium of Alcohol and Nicotine use; ICC, International Cannabis Consortium; ICD, International Classification of Diseases; iPSYCH, Lundbeck Foundation Initiative for Integrative Psychiatric Research; MVP, Million Veteran Program; PGC-ED, Eating Disorders Working Group of the Psychiatric Genomics Consortium; PGC-SUD, Substance Use Disorders Working Group of the Psychiatric Genomics Consortium; SNPs, single nucleotide polymorphisms.

<sup>&</sup>lt;sup>a</sup>Treated as a continuous phenotype.

<sup>&</sup>lt;sup>b</sup>Treated as an ordinal phenotype.

In Lui et al (2019), the phenotype is labeled as "smoking cessation." It was renamed as "current smoking" to reflect the coding scheme and for ease in comparing across all smoking phenotypes.

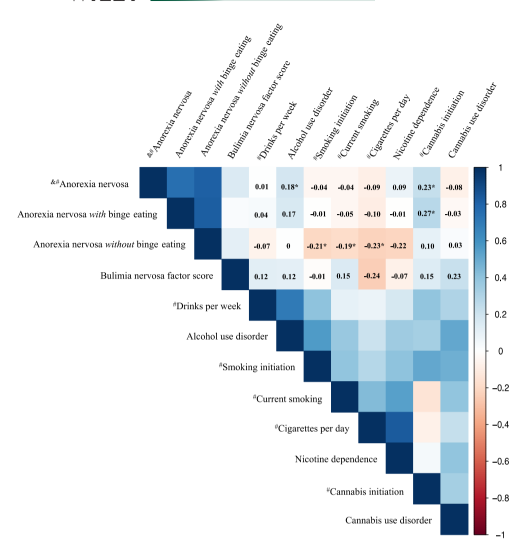


FIGURE 1 Genetic correlations between eating disorder subtypes and substance-use-related phenotypes. *Note.* # indicates known or potential sample overlap with UK Biobank, and & indicates known sample overlap with iPSYCH. Starred values denote significant genetic correlations after correcting for multiple comparisons using False Discovery Rate (n tests = 66; q < 0.05)

nonfunctional categories (corrected  $\rho_g$  = 0.01; corrected r = 0.27; corrected q = 0.004; Table S4).

Conversely, for smoking phenotypes, significant correlations were only observed for the AN *without* binge eating subtype. Smoking initiation ( $r_g = -0.21$ , SE = 0.06, q = 0.0006), current smoking (referred to as smoking cessation in Liu et al<sup>20</sup>)\* ( $r_g = -0.19$ , SE = 0.08, q = 0.03), and cigarettes per day ( $r_g = -0.23$ , SE = 0.07, q = 0.003) were significantly and negatively associated with AN *without* binge eating. Although the correlation between nicotine dependence and AN *without* binge eating was in the same direction as the other smoking phenotypes, it was not significant ( $r_g = -0.22$ , SE = 0.12, q = 0.14). The  $r_g$ s for AN diagnosis and each of the three nondiagnostic smoking traits versus AN *without* binge eating and these same smoking traits all differed significantly from each other (z-scores ranged from -3.22 to -2.11; P values  $\leq 0.04$ ). The genetic covariance estimate between AN *without* binge eating and smoking initiation was only significant in the nonfunctional category (corrected  $p_g = -0.01$ ; corrected r = 0.00).

-0.17; corrected q=0.007; Table S5). For AN without binge eating and current smoking, the genetic covariance estimate was significant in both functional (corrected  $\rho_g=-0.01$ ; corrected r=-0.32; corrected q=0.01) and nonfunctional categories (corrected  $\rho_g=-0.01$ ; corrected r=-0.21; corrected q=0.03; Table S6). Finally, the genetic covariance estimate between AN without binge eating and cigarettes per day was only significant in the nonfunctional category (corrected  $\rho_g=-0.02$ ; corrected r=-0.35; corrected q=0.003; Table S7).

After conditioning the AN and AUD GWAS summary statistics for loci associated with major depressive disorder, the positive  $r_g$  between AN and AUD was attenuated ( $r_g = 0.07$ ; SE = 0.05, q = 0.125; Table S8) and significantly lower than the unadjusted  $r_g$  (z-score = 2.48, P = 0.01). In contrast, after conditioning the AN with binge eating and cannabis initiation GWAS for major depressive disorder, the resulting  $r_g$  was marginally smaller but remained significant after correction for multiple tests ( $r_g = 0.21$ , SE = 0.08, q = 0.016). After conditioning for the major depressive disorder GWAS,  $r_g$ s between AN without binge eating and smoking initiation, current smoking, and cigarettes per day remained significant and modestly increased in magnitude ( $r_g$ s = -0.27 to -0.31; SEs = 0.05 to 0.09; qs < 0.0009). All  $r_g$ s

<sup>\*</sup>In Liu et al (2019), the phenotype is noted as "smoking cessation," where current smokers were coded as 2 and former smokers were coded as 1. Because the comparison group is "current smokers," we have renamed this phenotype as "current smoking" for clarification and ease of interpretation across all smoking phenotypes.

remained significant after conditioning the AN and substance-use-related phenotypes for schizophrenia ( $r_g$ s = -0.20 to 0.27; SEs = 0.04 to 0.08; qs < 0.03; Table S9).

## 4 | DISCUSSION

Using existing GWAS data, we investigated genetic associations between liabilities to four eating disorder- and eight substance-use-related phenotypes spanning initiation and typical use to substance use disorder. We found differential patterns of association between AN with and without binge eating and substance-use-related traits, which may point toward substance-specific genetic relationships. Additionally, there may be some degree of symptom overlap contributing to these associations.

Three main patterns emerged. First, in line with prior twin studies, we observed a positive genetic correlation  $(r_g)$  between problem alcohol use (ie, AUD) and AN diagnosis. Second, we observed positive, significant r<sub>o</sub>s between cannabis initiation and AN diagnosis, as well as cannabis initiation and the AN with binge eating subtype. This is a novel finding not previously examined in twin research. The positive genetic associations suggest that some genetic loci may be influencing these traits in the same direction. Finally, negative  $r_{o}$ s emerged between the three nondiagnostic smoking phenotypes and AN without binge eating but not with the other three eating disorder phenotypes. These negative  $r_g$ s indicate that some of the loci influencing liability to these eating disorder and smoking phenotypes might be shared but are affecting the liability to these traits in opposite directions. Indeed,  $r_{\sigma}$ s cannot identify specific loci or underlying mechanisms that contribute to the shared risk. Nevertheless, the results provide initial evidence for differential genetic associations between the liability to varying eating disorder- and substance-use-related phenotypes.

Based on findings from twin studies, we hypothesized that (a) the strongest SNP-based  $r_g$  would be between eating disorder phenotypes that have binge eating as a core symptom and alcohol use phenotypes and (b) a significant positive  $r_g$  between eating disorder phenotypes with binge eating as a key symptom and AUD would emerge. In line with these hypotheses, we found a significant genetic association between AUD and AN diagnosis but not between typical alcohol consumption (ie, drinks per week) and AN. No twin study has examined genetic associations between AN and alcohol-use-related phenotypes, and previous studies<sup>21,26</sup> using LDSC have not reported significant  $r_g$ s between these traits. That we found a significant association most likely reflects the larger AN sample size in our study (from 3495 cases and 10 982 controls to 16 992 cases and 55 525 controls), as well as combining two large existing GWAS of AUD, emphasizing the importance of increasing sample sizes for GWAS.

Importantly, the  $r_g s$  between the eating disorder- and substance-use-related phenotypes were robust to conditioning on schizophrenia loci. However, the  $r_g$  between AN and AUD was not robust to the adjustment for major depressive disorder-associated variants. Major depressive disorder is among the most prominent comorbidities in

individuals with AN and AUD,  $^{40}$  and GWAS for both traits document strong  $r_g$ s between major depressive disorder and these disorders.  $^{19,21,26}$  Our results indicate that the three disorders share genetic underpinnings. We cannot discount the possibility of a genetic relationship between AN and AUD that is distinct from major depressive disorder; however, much larger sample sizes may be required to detect such an association.

Intriguingly, although we did not detect a significant  $r_g$  for AN with binge eating with AUD, the point estimate for the  $r_g$  between AUD and AN with binge eating was similar to that for AUD and AN diagnosis (0.17 vs 0.18, respectively) and higher than that for AUD and AN without binge eating (0.01). Sample sizes for these AN subtypes were smaller than for AN diagnosis; however, the two subtypes included approximately equal numbers of cases and controls. Indeed, binge eating was assessed in such a way that we were unable to tease apart purging behaviors, and AN diagnosis is heterogenous even within subtypes. Therefore, binge eating may be one plausible key component of the observed genetic association. For example, binge eating has been shown to activate brain reward circuitry in a similar manner to substances, 41,42 and administration of naltrexone, an opioid antagonist approved by the US Food and Drug Administration for the treatment of AUD. 43 has been shown to reduce the frequency of binge eating episodes among individuals with an eating disorder. 44,45 We did not detect a significant  $r_g$  with the bulimia nervosa factor score, although that GWAS was relatively underpowered. Thus, our findings highlight the importance of expanding GWAS to include bulimia nervosa and binge-eating disorder, where a core symptom of both disorders is binge eating, to elucidate whether binge eating is a critical eating disorder symptom in the comorbidity with AUD and to home in on relevant shared mechanisms.

The significant genetic associations between cannabis initiation and AN are novel, yet consistent with the negative genetic association between cannabis use and body mass index, and with observational<sup>25</sup> and experimental<sup>46,47</sup> studies regarding the role of endocannabinoids in appetite regulation, energy expenditure, stress, and reward. One of the principal psychoactive agents of cannabis, delta-9-tetrahydrocannabinol (THC), a partial agonist of the endogenous cannabinoid 1 (CB1) receptor, is presumed to be orexigenic and may acutely increase appetite and food intake, contributing to its potential role as an appetite stimulant in patients with an anorexia or cachexia syndrome<sup>48</sup> due to a disease (eg, HIV or AIDS) or in response to treatment (eg, chemotherapy). An antagonist of the CB1 receptor was previously tested as a highly promising anti-obesity medication (Rimonabant, SR141716), which is particularly relevant because some genes may influence AN and obesity in opposite directions.<sup>21</sup> Further, the endocannabinoid anandamide has been shown to be elevated in individuals with acute AN,49 indicating disruption in food-related reward and eating behavior regulation. Animal and human studies have also provided initial evidence for the therapeutic effectiveness of cannabinoid agonists in treating eating disorders. 50,51 It is also likely that individuals with high genetic liability to AN are less likely to experiment with a substance that has a documented hyperphagia component. Thus, there is evidence of a complex

biological relationship between cannabis use and eating disorders, as well as body mass index.

Finally, the significant negative  $r_g$ s between three tobaccosmoking phenotypes-smoking initiation, current smoking, and cigarettes per day-and AN without binge eating are intriguing, suggesting that AN without binge eating and tobacco-smoking behaviors are alternate expressions of shared mechanisms. Phenotypic studies are inconsistent about the association between the restricting subtype of AN and smoking. Some studies suggest that individuals with restricting AN have a higher prevalence of various smoking phenotypes than controls,<sup>5</sup> whereas other studies indicate no significant difference between the two groups.<sup>6</sup> A recent metaanalysis did not find differences in the odds of lifetime smoking between individuals with AN and healthy controls,3 yet the authors did not assess differences by AN subtype. Individuals with AN may smoke as a way to control or lose weight, 52 and temporary weight gain does occur with smoking cessation.<sup>53</sup> However, a positive phenotypic correlation need not be accompanied by a  $r_g$  in the same direction (or genetic contributors to the phenotypic association at all). Still, there is plausible support for the negative  $r_g$ . Although not significant, a negative  $r_g$  between smoking and AN has been reported. 18,21 Notably, our study includes individuals from these earlier reports and extends findings by including larger sample sizes for both AN and smoking phenotypes. Unfortunately, there are no twin studies of AN or AN-like traits and smoking with which to compare findings.

One explanation for the negative genetic association is that it is due to a third, underlying variable influencing both AN without binge eating and smoking. We tested for the potential role of variants associated with major depressive disorder and schizophrenia and found the  $r_g$ s to be robust to those adjustments. In the largest GWAS of smoking phenotypes, positive  $r_q$ s were also observed between smoking initiation and cigarettes per day with multiple cardiometabolic traits, including type 2 diabetes and fasting glucose.<sup>20</sup> These same metabolic traits were negatively genetically correlated with AN.  $^{21,54}$  Thus, the patterns of  $r_{\rm e}$ s might point to metabolic, rather than psychiatric, factors in influencing the apparent genetic association between smoking phenotypes and AN. However, the associations could also reflect adoption of unhealthy lifestyles that promote obesity and are correlated with smoking. In addition, the  $r_{\sigma}$ s between smoking and body mass index, as well as AN and body mass index, may reflect underlying disinhibitory pathways, as variants associated with body mass index show enrichment in the central nervous system.<sup>55</sup> The current approach is not designed to disentangle these putative etiological mechanisms, but our findings do encourage careful study of the specific relationships between eating and substance use disorders.

Substance use and substance use disorders are partially distinct, and although excessive substance use is a necessary component of substance use disorders, the latter is associated with psychological and physiological impairment related to excess use and aspects of loss of control over the behavior. Consistent with our findings for alcohol, accumulating evidence suggests that genetic liability to other

psychiatric traits (eg, schizophrenia) is strongly correlated with liability to substance use disorders (eg, AUD) but not substance use (eg, alcohol consumption). 19-21 Genetic liability to alcohol use has also been correlated with liabilities to psychiatric disorders (eg, major depressive disorder) in opposite directions depending on level of involvement. 19 However, we did not find similar elevations in  $r_{g}$ s when contrasting ever smoking and nicotine dependence nor comparing cannabis initiation to cannabis use disorder. It is possible that the lack of genetic overlap between AN and nicotine dependence, as well as AN and cannabis use disorder, is related to the relatively modest sample size of those discovery GWAS. A similar non-significant  $r_g$  was noted for AUD when the Walters et al<sup>26</sup> alcohol dependence GWAS was used as the sole source of summary statistics for problem drinking in the current study. Several other explanations for this divergence in findings exist. For instance, for tobacco, the highly addictive nature of nicotine may result in convergence in genomic effects on earlier and later stages of smoking (ie, a much larger proportion of those who ever smoke become dependent compared with the proportion of those who drink alcohol and develop AUD). For cannabis, given its lower addictive potential, we might have expected stronger associations with cannabis use disorder than with cannabis initiation. In addition to the considerably smaller sample size of the cannabis use disorder GWAS, the association with cannabis initiation could also be attributed to the small number of cohorts in that discovery GWAS that included individuals with a high likelihood of cannabis use disorder. It is also possible that the relationship between AN and cannabis use is distinct and that earlier but not later stages of cannabis use are genetically related to liability to AN. Future studies should consider the multistage nature of substance use and misuse when examining cross-trait correlations.

This is the largest and most comprehensive assessment of shared genetic risk between eating disorder- and substance-userelated phenotypes, using existing GWAS data from large cohorts (up to ~537 000 individuals per phenotype). We were able to separately assess approximate AN subtypes (ie, with binge eating vs without binge eating) to evaluate the extent to which binge eating, in the context of AN, may share genetic risk with substance-userelated phenotypes. Using these large datasets-many of which are publicly available-allows for the rapid development of scientific knowledge regarding the underlying etiology of psychiatric disorder and substance use comorbidity. Nevertheless, some limitations exist. First, sample sizes for the bulimia nervosa factor score and cannabis use disorder GWAS were relatively small compared with the other GWAS, resulting in large standard errors and low power. Second, we were unable to uniformly examine sex differences in these r<sub>g</sub>s. Because the prevalence of eating disorders is higher in women than men and the prevalence of substance use disorders is higher in men than women,40 it will be important to explore possible sex differences in genetic associations as the GWAS data become available. Notably, we previously did not find evidence for sex differences in the  $r_g$  between binge eating and problem alcohol use.<sup>56</sup> Third, even though we did not detect significant rgs for all pairs of traits, it is possible that local genetic associations exist for some of these trait

pairs. Such local correlations, for instance, in certain chromosomal regions but not others, particularly when in opposing directions (eg, a positive local correlation at one chromosomal location and a negative local correlation at another) might dilute the overall  $r_g$  estimate. Although such a systematic evaluation of each pair of traits is beyond the scope of this report, we did note some support for enrichment of the aggregated genetic covariance in both functional and nonfunctional genomic regions for several of the significant r<sub>g</sub>s. Finally, SNP coverage was limited in the earlier GWAS of the bulimia nervosa factor score because that study used older genotyping platforms and imputation panels that included fewer SNPs than current imputation panels. The Eating Disorders and Substance Use Disorders Working Groups of the Psychiatric Genomics Consortium are continuously adding samples and releasing data freezes with incrementally larger sample sizes, while collecting information on multiple substances (eg, opioids). In coming years, the statistical power is expected to increase for AN (including the with and without binge eating subtypes), bulimia nervosa, and bingeeating disorder, as well as AUD, nicotine dependence, and cannabis use disorder, from within and outside the Psychiatric Genomics Consortium. This will allow for a more refined assessment of specific eating disorder symptoms, including binge eating, in relation to substance-use-related phenotypes.

In conclusion, findings from this study suggest that the shared sources of variation in liabilities to eating disorder- and substanceuse-related phenotypes are not consistent across traits or levels of substance involvement, extending results from twin studies to a genome-wide SNP approach. Despite the typically high cooccurrence of alcohol, tobacco, and cannabis use and their genetic overlap.<sup>25</sup> the differential patterns seen between the eating disorder- and substance-use-related phenotypes highlight the uniqueness and complexity of their shared etiology. Potential clinical implications include watching for the emergence of symptoms of one disorder (eg, AN) while being treated for the other behavior (eg, AUD) and understanding that, for example, women with AN who use nicotine may not be able to quit successfully both because they are afraid of gaining weight and they have high genetic susceptibility for smoking via the shared genetic risk between AN and smoking-related traits. Additional research using contemporary genomic methods, such as cross-disorder association studies, could identify the specific loci contributing to this comorbidity. Future research that combines genome-wide data with measured environmental constructs, such as trauma,9 that may increase risk for this comorbidity could enhance the prediction, prevention, and treatment of co-occurring eating disorder- and substance-use-related traits.

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# Eating Disorders Working Group of the Psychiatric Genomics Consortium (PGC-ED)

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# Substance Use Disorders Working Group of the Psychiatric Genomics Consortium (PGC-SUD)

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#### **DATA ACCESS**

This manuscript was a joint collaboration between the Eating Disorders and Substance Use Disorders Working Groups of the Psychiatric Genomics Consortium. These data can be found at https://www.med.unc.edu/pgc/data-index/. Additional datasets included in this study were obtained multiple ways. We recieved summary statistics directly from the first author of the primary GWAS manuscript for the bulimia nervosa factor score (Australian Twin Registry), alcohol use disorder (Million Veteran Program), nicotine dependence (multiple samples), and cannabis initiation (International Cannabis Consortium and UK Biobank). Summary statistics for drinks per week, smoking initiation, smoking cessation, and cigarettes per day (GSCAN) were downloaded from https://conservancy.umn.edu/handle/11299/201564 on 7 March 2019. Summary statistics for cannabis use disorder (iPSYCH) were downloaded from https://ipsych.dk/forskning/downloads/ on 27 June 2019.

#### **CONFLICT OF INTERESTS**

The authors report the following potential competing interests. O. Andreassen received a speaker's honorarium from Lundbeck. G. Breen received grant funding and consultancy fees in preclinical genetics from Eli Lilly, consultancy fees from Otsuka, and has received honoraria from Illumina. C. Bulik served on Shire Scientific Advisory Boards, is a consultant for Idorsia, and receives author royalties from Pearson. D. Degortes served as a speaker and on advisory boards and has received consultancy fees for participation in research from various pharmaceutical industry companies including AstraZeneca, Boehringer, Bristol Myers Squibb, Eli Lilly, Genesis Pharma, GlaxoSmithKline, Janssen, Lundbeck, Organon, Sanofi, UniPharma, and Wyeth; he has received unrestricted grants from Lilly and AstraZeneca as director of the Sleep Research Unit of Eginition Hospital (National and Kapodistrian University of Athens, Greece). J. Hudson has received grant support from Shire and Sunovion and has received consulting fees from DiaMentis, Shire, and Sunovion. A. Kaplan is a member of the Shire Canadian Binge-Eating Disorder Advisory Board and was on the steering committee for the Shire B/educated Educational Symposium: 15 to 16 June 2018. J. Kennedy served as an unpaid member of the scientific advisory board of AssurexHealth Inc. M. Landén declares that, over the past 36 months, he has received lecture honoraria from Lundbeck and served as scientific consultant for EPID Research Oy. S. Scherer is a member of the scientific advisory board for Deep Genomics. P. Sullivan is on the Lundbeck advisory committee and is a Lundbeck grant recipient; he has served on the scientific advisory board for Pfizer, has received a consultation fee from Element Genomics and a speaker reimbursement fee from Roche. J. Treasure has received an honorarium for participation in an EAP meeting and has received royalties from several books from Routledge, Wiley, and Oxford University press. T. Werge has acted as a lecturer and scientific advisor to H. Lundbeck A/S. L. Bierut, A. Goate, J. Rice, J.-C. Wang, and the spouse of N. Saccone are listed as inventors on Issued US Patent 8080,371, "Markers for Addiction" covering the use of certain SNPs in determining the diagnosis, prognosis, and treatment of addiction. N. Wodarz has received funding from the German Research Foundation (DFG) and Federal Ministry of Education and Research Germany (BMBF); he has received speaker's honoraria and travel funds from Janssen-Cilag, Mundipharma, and Indivior. He took part in industrysponsored multicenter randomized trials by D&A Pharma and Lundbeck, M. Ridinger received compensation from Lundbeck Switzerland and Lundbeck institute for advisory boards and expert meetings and from Lundbeck and Lilly Suisse for workshops and presentations. K. Mann received speaker fees from Janssen-Cilag. H. Kranzler is a member of the American Society of Clinical Psychopharmacology's Alcohol Clinical Trials Initiative, which was sponsored in the past 3 years by AbbVie, Alkermes, Amygdala Neurosciences, Arbor Pharmaceuticals, Ethypharm, Indivior, Lilly, Lundbeck, Otsuka, and Pfizer. H. Kranzler and J. Gelernter are named as inventors on PCT patent application #15/878.640, entitled "Genotype-guided dosing of opioid agonists," filed 24 January 2018. J. MacKillop is a principal in BEAM Diagnostics, Inc. D.-S. Choi is a scientific advisory member of Peptron Inc. M. Frve has received grant support from Assurex Health, Mayo Foundation, Myriad, National Institute on Alcohol Abuse and Alcoholism. National Institute of Mental Health, and Pfizer; he has been a consultant for Intra-Cellular Therapies, Inc., Janssen, Mitsubishi Tanabe Pharma Corporation, Myriad, Neuralstem Inc., Otsuka American Pharmaceutical, Sunovion, and Teva Pharmaceuticals. H. de Wit has received support from Insys Therapeutics and Indivior for studies unrelated to this project, and she has consulted for Marinus and Jazz Pharmaceuticals, also unrelated to this project. T. Wall has previously received funds from ABMRF. J. Nurnberger is an investigator for Janssen. M. Nöthen has received honoraria from the Lundbeck Foundation and the Robert Bosch Stiftung for membership on advisory boards. N. Scherbaum received honoraria for several activities (advisory boards, lectures, and manuscripts) by the factories Abbvie, Hexal, Janssen-Cilag, MSD, Medice, Mundipharma, Reckitt-Benckiser/Indivior, and Sanofi-Aventis. W. Gäbel has received symposia support from Janssen-Cilag GmbH, Neuss, Lilly Deutschland GmbH, Bad Homburg, and Servier, Munich and is a member of the Faculty of the Lundbeck International Neuroscience Foundation (LINF), Denmark. J. Kaprio has provided consultations on nicotine dependence for Pfizer (Finland) 2012 to 2015. In the past 3 years, L. Degenhardt has received investigator-initiated untied educational grants for studies of opioid medications in Australia from Indivior, Mundipharma, and Segirus. B. Neale is a member of the scientific advisory board for Deep Genomics and has consulted for Camp4 Therapeutics Corporation, Merck & Co., and Avanir Pharmaceuticals, Inc. A. Agrawal previously received peer-reviewed funding and travel reimbursement from ABMRF for

unrelated research. All other authors have no conflicts of interest, relevant to the contents of this paper, to disclose.

#### **AUTHORS CONTRIBUTION**

MM-C, CB, and AA were responsible for the study concept and design. MM-C, ECJ, and Y-LD performed the statistical analyses, and JC, RW, and ZY assisted with the data analysis. MM-C, ECJ, Y-LD, JC, LT, RW, ZY, JB, CH, JK, HE, CB, and AA assisted with interpretation of findings. TDW facilitated access to and interpretation of the summary statistics for the bulimia nervosa factor score. HK, JG, and HZ facilitated access to and interpretation of the Million Veteran Program summary statistics for AUD. DH facilitated access to and interpretation of the summary statistics for nicotine dependence. MM-C, ECJ, LT, CB, and AA drafted the manuscript. All remaining authors provided data for this study and consulted on the analytic plan. All authors critically reviewed the content and approved the final version for publication.

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#### SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of this article.

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