The etiology of DSM-5 alcohol use disorder: Evidence of shared and non-shared additive genetic effects

Rohan H.C. Palmer a,⁎, Leslie A. Brick b,c, Yi-Ling Chou d, Arpana Agrawal d, John E. McGeary b,c,e, Andrew C. Heath d, Laura Bierut d, Matthew C. Keller f, Eric Johnson g, Sarah M. Hartz d, Marc A. Schuckit h, Valerie S. Knopik i

a Behavioral Genetics of Addiction Laboratory, Department of Psychology, Emory University, USA
b Department of Psychiatry and Human Behavior, Brown University, USA
c Division of Behavior Genetics, Department of Psychiatry, Rhode Island Hospital, USA
d Washington University in St. Louis, St. Louis, Missouri, USA
e Providence Veterans Affairs Medical Center, USA
f Department of Psychology and Neuroscience, University of Colorado at Boulder, USA
g RTI International, USA
h University of California, San Diego, USA
i Department of Human Development and Family Studies, Purdue University, USA

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ABSTRACT

Background: Alcoholism is a multifactorial disorder influenced by multiple gene loci, each with small effect. Studies suggest shared genetic influences across DSM-IV alcohol dependence symptoms, but shared effects across DSM-5 alcohol use disorder remains unknown. We aimed to test the assumption of genetic homogeneity across the 11 criteria of DSM-5 alcohol use disorder (AUD).

Methods: Data from 2596 alcohol using individuals of European ancestry from the Study of Addiction: Genetics and Environment were used to examine the genomewide SNP-heritability (h²SNP) and SNP-covariance (rGSNP) between 11 DSM-5 AUD symptoms. Phenotypic relationships between symptoms were examined to confirm an underlying liability of AUD and the SNP-heritability of the observed latent trait and the co-heritability among AUD symptoms was assessed using Genomic-Relatedness-Matrix-Restricted-Maximum-Likelihood. Genetic covariance among symptoms was examined using factor analysis.

Results: Phenotypic relationships confirmed a unidimensional underlying liability to AUD. Factor and parallel analyses of the observed genetic variance/covariance provided evidence of genetic homogeneity. Additive genetic effects on DSM-5 AUD symptoms varied from 0.10 to 0.37 and largely overlapped (rG-SNP across symptoms ranged from 0.49 - 0.92). The additive genetic effect on the DSM-5 AUD factor was 0.36, 0.14 for DSM-5 AUD diagnosis, and was 0.22 for DSM-5 AUD severity.

Conclusions: Common genetic variants influence DSM-5 AUD symptoms. Despite evidence for a common AUD factor, the evidence of only partially overlapping genetic effects across AUD symptoms further substantiates the need to simultaneously model common and symptom-specific genetic effects in molecular genetic studies in order to best characterize the genetic liability.

1. Introduction

Until recently, alcohol abuse and dependence, as described by the Diagnostic and Statistical Manual of Mental Disorders (DSM IV; (American Psychiatric Association, 2000)), were the most studied problematic outcomes for the clinical consequences of alcohol consumption. As diagnostic measures, alcohol abuse or dependence were restricted to classifying individuals as “affected” or “unaffected” with little sensitivity for underlying profiles of endorsement of the 11 symptoms (7 for dependence and 4 for abuse) that were being used to describe alcohol-related behavior. Hence, especially when used as research outcomes, there was significant concern about DSM-IV diagnoses...
of abuse or dependence to (1) reflect individual differences with respect to the underlying constructs believed to be represented by the 11 symptoms that research showed is indicative of a single continuum (Hasin and Bresler, 2009; NRC, 2011), and (2) reflect differences in severity across all of the addiction domains (i.e., (a) a compulsion to seek and/or take alcohol, (b) loss of control over alcohol consumption, and (c) emergence of a negative emotional state) captured by the symptoms. Not surprising, recent studies and recommendations which arose out of the Substance Related Disorders Working Group of the DSM-5 taskforce suggested that diagnosis of an alcohol use disorder (AUD) be based on 11 symptoms derived from the integration of the DSM-IV dependence symptoms with three of the DSM-IV abuse criteria (i.e., ‘recurrent legal problems’) and an alcohol craving criterion. Notably, a continuum should be used to describe AUDs (i.e., unaffected = endorsing 0–1 out of 11 symptoms; mild = endorsing 2–3 out of 11 symptoms; moderate = endorsing 4–5 symptoms out of 11 symptoms, and severe = endorsing 6 or more out of 11 symptoms).

As part of the debate on the utility and suitability of the dichotomous measure of alcohol dependence (AD), as opposed to a continuous measure of alcohol problem severity in genetic research, we recently examined the assumption of genetic homogeneity across all DSM-IV dependence symptoms using genomewide SNP data (Palmer et al., 2015b). Validation of the assumption across the seven symptoms affirmed the utility of a factor score across the indices as much of the observed genetic variance was shared across the comorbid items, suggesting common genetic factors underlie the addiction state (Koob et al., 2014), which is reflected in behavioral symptoms included in the addiction state (American Psychiatric Association, 2013). The results also indicated that effects observed upon a latent continuum of AD risk (as indicated by DSM-IV dependence symptoms) may not be truly reflective of the entire liability continuum, as there also exists symptom-specific genetic variance that may be imparted by the study of multiple factors (as previously suggested using a multivariate twin study approach (Kendler et al., 2012)). This latter point was recently reflected in a report by Hart and colleagues which showed variation in the association between common genetic variants within the alcohol dehydrogenase gene (ADH1B) and each of the diagnostic symptoms of AD (Hart et al., 2016). In perspective, Hart and colleagues were able to determine that previously observed AD genomewide association study (GWAS) associations (Gelernter et al., 2014), for example, in their subjects of African ancestry (i.e., rs2066702 with AD), were primarily driven by signals specific to phenotypic variation in the symptoms ‘Tolerance’ and ‘Much time spent using/recovering from the effects of alcohol’. This observation is important as phenotype-genotypic associations from GWAS are used to inform gene function studies in tissue/cell culture and/or model organisms.

Altogether, our previous study of DSM-IV symptoms and these recent molecular studies of DSM-IV and DSM-5 AUD underscore the need to characterize the multivariate genetic architecture of DSM-5 AUD symptoms. The present paper uses subjects of European ancestry from the Study of Addiction: Genetics and Environment Consortium to characterize the genetic architecture of the 11 DSM-5 AUD symptoms. It builds upon our previous report by comparing several models that test the assumption of genetic homogeneity across AUD symptoms. The goals of this study were to:

1. Examine the genomewide additive genetic contribution to DSM-5 AUD symptoms (i.e., former DSM-IV abuse and dependence symptoms along with alcohol craving), and
2. Determine the most parsimonious model of the additive genetic covariance across DSM-5 AUD symptoms.

2. Materials and methods

2.1. Sample

Data were drawn from the Study of Addiction: Genetics and Environment (SAGE) (Bierut et al., 2010). Analyses focused on 2596 unrelated individuals (44% male; mean age = 38.58 years [standard deviation (SD) = 9.80]) of European ancestry, which was confirmed using principal component analysis. All subjects were no more related than second cousins (Palmer et al., 2015a). Additional details on SAGE are available at http://www.ncbi.nlm.nih.gov/projects/gap/cgi-bin/study.cgi?study_id=phs000092.v1.p1.

2.2. Phenotype

The dependent variables of interest for the current study were DSM-5 AUD symptoms (coded as present or absent) that were approximated from the Semi-Structured Assessment for the Genetics of Alcoholism (SSAGA) (Bucholz et al., 1994; Hesselbrock et al., 1999). Specifically, DSM-IV alcohol abuse and dependence symptoms were extracted from the SSAGA interview and combined with a separate measure of alcohol craving that was also included in the assessment. Craving was defined in the SSAGA as endorsement of the item ‘In situations where you couldn’t drink, did you ever have such a strong desire for it that you couldn’t think of anything else’; respondents who answered yes received a score of 1; ‘No’ was coded as 0; non-users received a missing value (Agrawal et al., 2013). All responses were limited to individuals with a history of exposure to alcohol (and possibly other substances).

2.3. Genotyping and quality control

Subjects within SAGE were genotyped using the ILLUMINA Human 1 M platform. Quality control of the sample included: (1) removal of non-autosomal SNPs, (2) removal of markers with an allele frequency < 1%, (3) exclusion of markers with a call rate less than 98%, and (4) removal of SNPs that show evidence of deviation from Hardy–Weinberg Equilibrium (HWE; p-value < 0.0001) to minimize any possible bias due to assortative mating (Agrawal et al., 2006; Grant et al., 2007). A total of 796,125 autosomal SNPs were carried forward in the analyses. These same SNPs were also used to conduct the aforementioned selection of distantly related EA individuals from the entire set of SAGE participants (N = 4121) (i.e., using the software package: Genomewide Complex Trait Analysis (GCTA))(Yang et al., 2010; Lee et al., 2011; Yang et al., 2011).

2.4. Statistical analysis

The EA sample (N = 2596) was used in all parts of the analytical framework, which included (1) development of a phenotypic factor comprised of shared variance among DSM-5 items using randomly selected individuals to create two halves of the sample for exploratory and confirmatory models (conducted in Mplus version 7) (Muthén and Muthén (1998-2012))Muthén and Muthén, - , 2012Muthén and Muthén (1998-2012)), (2) Genomic-relatedness-matrix restricted maximum likelihood (GREML; implemented in GCTA) analysis of the individual symptoms, AUD factor score (standardized [mean = 0, standard deviation = 1]), AUD diagnosis (i.e., 0 = 0–1 symptoms, 1 = 2–11 symptoms), and log-transformed DSM-5 AUD diagnosis severity (i.e., Ln (1 + DSM-5 AUD diagnosis [0 = 0–1 symptoms, 1 = mild [2–3 symptoms], 2 = moderate [4–5 symptoms], 3 = severe [7 + symptoms])).
2.4.1. Estimation of additive genetic effects

GREML was used to determine the SNP-heritability ($h^2_{SNP}$) of DSM-5 AUD factor score, severity (i.e., mild, moderate, severe), diagnosis (i.e., control vs. case [i.e., 2+ symptoms]), and individual symptoms. This approach was implemented using GCTA. GREML utilizes a genetic relationship matrix to decompose phenotypic variance into genetic effects captured by the common SNPs and error variance. The SNP-heritability estimates were transformed on the liability scale to account for distributional differences in prevalence of AUD and endorsement of AUD symptoms observed in this case/control study versus the general population (i.e. the proportion of cases in this study is higher than what is seen in the population). Lifetime population prevalence estimates that were used to transform the SNP-heritability and co-heritability estimates were calculated for DSM-5 AUD diagnosis and individual symptoms from the National Epidemiologic Survey on Alcohol and Related Conditions (NESARC, wave 1; N = 43,093) (Hasin and Grant, 2015) and, for craving from the National Longitudinal Alcohol Epidemiologic Survey (NLAES) (Grant et al., 2003). Craving in NLAES was defined by endorsement of at least one of two possible items: ‘Want to drink so badly that you couldn’t think of anything else’ and ‘Feel a very strong desire or urge to drink’. Prevalence was calculated for individuals in NESARC and NLAES who (1) self-reported non-Hispanic White ethnicity, (2) were aged 18–79 years, and (3) reported lifetime exposure to alcohol (see Table 2). All analyses controlled for gender, age, and the first five ancestral principal components.

2.4.2. Estimation of the covariance explained by SNPs

Two multivariate approaches were used to determine whether the same genetic factors contribute to the phenotypic correlation between AUD symptoms: (1) The Common Pathway Model (CPM) and (2) Exploratory Genetic Factor Analysis (EGFA). In addition, the EGFA was followed up with three confirmatory factor analyses to determine the most parsimonious genetic architecture across the criteria. Three multivariate models were tested: (a) a common genetic factor model, (b) a 2-factor model where factor-1 was indicated by the three former DSM-IV abuse symptoms and craving and factor-2 was indicated by the seven former DSM-IV dependence items, and (c) a 2-genetic factor model in which craving was allowed to cross-load across the factors. The CFA models were compared using the Akaike Information Criteria (AIC) (Akaike, 1973).

In the CPM approach, a latent variable representing the shared variance across all symptoms was decomposed into genetic and error variance in two steps. First, an exploratory factor model (EFA) was fitted to a random selected half of the sample to determine the phenotypic factor structure of AUD; this model was then confirmed using confirmatory factor analysis (CFA) of the remaining half of the sample. AUD factor scores were extracted from the full sample and used in the analyses described above. In the EGFA approach, which represents a multivariate extension of GREML, a factor analysis was conducted on the 11 × 11 variance/covariance matrix of inter-criterion bivariate SNP heritabilities through a series of steps. First, GREML was used to estimate bivariate SNP genetic covariance estimates across each pair of criteria. Next, these estimates were used to construct an 11 × 11 genetic variance/covariance matrix. Because covariance matrices constructed from bivariate estimates may not be positive definite, we determined the nearest positive definite variance/covariance matrix using the Higham algorithm (Higham, 2002) within the nearPD package in R, version 3.4.0 (Team, 2017). Finally, we conducted factor analysis of the variance/covariance matrix to determine the factor structure of the multivariate genetic relationship between AUD symptoms. To determine the number of genetic factors, we employed Parallel Analysis implemented in R with the nFactors package. This approach has been shown to outperform other methods under a variety of conditions (Ledesma and Valero-Mora, 2007). A factor was retained if the eigenvalue of the genetic variance/covariance matrix was greater than the 95th percentile of the distribution of eigenvalues derived from random data (generated with 1000 iterations). All analyses for the CPM approach were conducted in Mplus and all analyses for the EGFA approach were conducted using Mplus and the R version 3.4.0 packages.
3. Results

3.1. Symptom levels and phenotypic covariance in SAGE

The prevalence of endorsement for each AUD symptom and the tetrachoric correlations among all items are presented in Table 1. Approximately 66.49% (n = 1716) of participants met the diagnostic symptoms for DSM-5 AUD. Phenotypic tetrachoric correlations were generally high (strong rG-SNP > 0.60), suggesting shared genetic relationships between symptoms as arising from an unobserved latent trait. Common SNPs explained 36% (standard error [SE] = 0.13, p = 0.002) of the variation in the AUD factor score. However, it is important to recognize that several of these estimates were inflated in instances where the heritability of at least one of the symptoms was non-significant (i.e., only a small proportion of the phenotypic variance in the DSM-5 criterion is explained by genetic variation). Analysis of the 11 × 11 genetic variance/covariance matrix suggested a single genetic factor parsimoniously describes much of the shared genetic variance across the 11 criteria (see Supplementary Table S1 for the genetic variance/covariance matrix). Parallel analysis indicated that the first eigenvalue derived from this matrix exceeded the 95th percentile of the distribution of eigenvalues derived from random data (see Fig. 1). Genetic factor loadings for AUD symptoms were high (>0.60) and the total genetic variance of each criteria attributable to the factor ranged from 38% for ‘craving’ to 89% for ‘failure to fulfill major roles’ (see Table 4 for a summary of factor loadings and percent variance explained in the EGFA). Our analysis of competing models of additive genetic effects on AUD indicated that the model containing a single genetic factor provided the best fit to the data (χ² = 49905.53, degrees of freedom (df) = 44, AIC = 49817.53). On the contrary, the two-factor genetic model that allowed for a correlation between the genetic factors was less parsimonious (compared to the one-factor model: ΔAIC = 4046.55), but estimated the correlation between the abuse and dependence factors at 1.00 [95% confidence interval=0.99, 1.00]. We also examined a bi-genetic factor model that allowed craving to cross-load across the factors; results were similar and this adjusted model fit the single latent factor > 0.84. Excellent model fit (RMSEA < 0.05, CFI/TLI > 0.95) supports the CPM that describes the phenotypic relationships between symptoms as arising from an unobserved latent trait. Common SNPs explained 36% (standard error [SE] = 0.13, p = 0.002) of the variation in the AUD factor score.

3.2. Univariate additive genetic effects on DSM-5 AUD diagnosis and symptoms

Common SNPs explained 14% (standard error [SE] = 0.21, p = 0.24 of the variation in DSM-5 AUD diagnosis and 22% (SE = 0.13, p = 0.04) of the variation in AUD severity (i.e., In-transformed DSM-5 AUD categories). Across the 11 AUD symptoms, SNP-heritability estimates varied from 13% (Great time spent using/recovering) to 39% (Using over a longer period than was intended), with five of the 11 items reaching significance (p < 0.10) and four items reaching nominal significance (p < 0.05) and two criteria: tolerance and ‘Taking alcohol in larger amounts or over a longer period than was intended’.

3.3. Analysis of the genetic covariance across DSM-5 AUD symptoms

3.3.1. CPM approach

Exploratory and confirmatory analyses of phenotypic data revealed a single latent variable (AUD factor; see Table 3). All items loaded on the single latent factor > 0.84. Excellent model fit (RMSEA < 0.05, CFI/TLI > 0.95) supports the CPM that describes the phenotypic relationships between symptoms as arising from an unobserved latent trait. Common SNPs explained 36% (standard error [SE] = 0.13, p = 0.002) of the variation in the AUD factor score.

3.3.2. EGFA approach

Across AUD symptoms, the pattern of inter-symptom SNP correlations was generally high (strong rG-SNP > 0.60), suggesting shared genetic variance across symptoms (see right side of Table 2). However it is important to recognize that several of these estimates were inflated in instances where the heritability of at least one of the symptoms was non-significant (i.e., only a small proportion of the phenotypic variance in the DSM-5 criterion is explained by genetic variation). Analysis of the 11 × 11 genetic variance/covariance matrix suggested a single genetic factor parsimoniously describes much of the shared genetic variance across the 11 criteria (see Supplementary Table S1 for the genetic variance/covariance matrix). Parallel analysis indicated that the first eigenvalue derived from this matrix exceeded the 95th percentile of the distribution of eigenvalues derived from random data (see Fig. 1). Genetic factor loadings for AUD symptoms were high (> 0.60) and the total genetic variance of each criteria attributable to the factor ranged from 38% for ‘craving’ to 89% for ‘failure to fulfill major roles’ (see Table 4 for a summary of factor loadings and percent variance explained in the EGFA). Our analysis of competing models of additive genetic effects on AUD indicated that the model containing a single genetic factor provided the best fit to the data (χ² = 49905.53, degrees of freedom (df) = 44, AIC = 49817.53). On the contrary, the two-factor genetic model that allowed for a correlation between the genetic factors was less parsimonious (compared to the one-factor model: ΔAIC = 4046.55), but estimated the correlation between the abuse and dependence factors at 1.00 [95% confidence interval=0.99, 1.00]. We also examined a bi-genetic factor model that allowed craving to cross-load across the factors; results were similar and this adjusted model fit
the data slightly worse (compared to the one-factor model: ΔAIC = 4048.55). Overall, both the EGFA and CFA suggest shared additive genetic effects across symptoms of DSM-5 AUD.

4. Discussion

This study examined the expanded definition of diagnostic criteria contributing to AUD as defined by the American Psychiatric Association’s DSM-5. Additive genetic effects are partially shared across DSM-5 symptoms of AUD, with genetic correlations > 0.80 for several criteria. However, correlations across some criteria were as low as 0.21 (e.g., craving and time spent), suggesting the possibility of a violation of the assumption of genetic homogeneity underlying the AUD phenotype, but this was not a common occurrence (i.e., percentage of correlations > 0.3 = 98%, > 0.6 = 78%, and > 0.8 = 42%). Notably, while the standard errors for some of these correlation estimates was fairly large (e.g., rG = 0.21 (SE = 0.73) for the association between craving and ‘A great deal of time spent to obtain/use/recover from alcohol’), other correlations were more precise (e.g., rG = 1.00 (SE = 0.20) for ‘Recurrent use resulting in failure to fulfill major roles’ associated with ‘Given up or cut back on important activities in order to drink’), and thus, the hypothesis of underlying genetic homogeneity cannot be fully rejected. However, it is possible that the ability to localize genetic loci for AUDs is likely to be reduced when using scoring methods that ignore the fact that symptoms are influenced by shared and non-shared genetic factors (i.e., just as there are a multitude of symptom profiles that lead to an AUD diagnosis, the respective genetic risk profiles for these various symptom profiles may vary accordingly).

The incorporation of previous DSM-IV abuse symptoms and craving for alcohol enhanced the definition of problematic alcohol use. Estimated effects of genetic variation on the newly added alcohol symptoms ranged from 0.10-0.37. Notably, the SNP-heritability of DSM-5 AUD factor was similar to what was previously reported for DSM-IV alcohol dependence as a factor score (Palmer et al., 2015b; Brick et al., 2017). Moreover, our examinations of the genetic influences on the 11 symptoms supports the underlying assumptions of (1) a single underlying dimension of risk that is captured by the symptoms, and (2) common genetic pathways that contribute to ‘craving’, ‘using longer than intended’, ‘withdrawal’, and the other symptoms. These findings align with previous examinations of alcohol symptoms in genetically informed samples, which suggested a single underlying latent trait that is polygenic in nature. It was for this reason that we opted to use Genomic Restricted maximum likelihood (GREML) to understand the relationship among the 11 AUD symptoms because our sample sizes precluded the use of genomewide association analysis which would have resulted in biased SNP-estimates that reflect only a small portion

Note: EFA = Exploratory factor analysis; CFA = Confirmatory factor analysis; df = degrees of freedom; RMSEA = Root Mean Square Error Of Approximation; CFI = Comparative Fit Index; TLI = Tucker Lewis Index; RMSR = Root Mean Square Residual

![Fig. 1. Parallel analysis of 11 × 11 genetic covariance matrix for DSM-5 AUD symptoms. Observed eigenvalues (solid line) are compared to 95 percentile of the eigenvalue distribution (dashed line [with standard error]) derived from 1000 randomly generated datasets. All factors left of where the solid lines first intersects with the dashed line are retained and their effects described in Table 4.](image-url)
of the phenotypic heritability (i.e., missing heritability; (Manolio et al., 2009)). In the current analysis, we were interested in quantifying the heritability and co-heritability due to common variation. In GREML, there is less emphasis on detecting the small effects of the common variants, but instead more emphasis on aggregated effects. Our examination of alternative factorial configurations of the criteria (i.e., correlated abuse and dependence genetic factors) provides novel evidence supporting the assumption that the genetic architecture across AUD symptoms is largely shared. Indeed, as the data suggest, GWAS aimed at the factors of tolerance, loss of control and withdrawal (Kendler et al., 2011) may yield loci distinct from those identified using a unidimensional factor score or diagnosis, because of limited power to detect such specificity and also because of AUD-symptom-specific genetic variance. As such, future works should consider analyzing AUD measures in their various forms.

Using these approaches, for the first time we report on the genomewide SNP-heritability of alcohol craving. Similar to our earlier report using a larger, but ancestrally mixed SAGE sample (Agrawal et al., 2013), 21% percent of the total sample endorsed alcohol craving. The univariate SNP-heritability of 0.24 (SE = 0.21; post-hoc power = 0.57) for craving did not meet our criteria for statistical significance, however, the high loading (> 0.85) of the craving item on the latent AUD factor, which had a SNP-heritability of 0.36 (standard error [SE] = 0.13), suggests genetic effects on craving. Notably, our analysis of the genetic effects across symptoms suggested some differential effects of genomewide SNPs across AUD symptoms, but craving was least explained by the common genetic factors. A review of the alcohol literature identified two studies supporting the role of variation across the alpha-synuclein gene (SNCA) and alcohol craving. α-synuclein has been shown to play a role in dopamine functioning across several regions of the brain (e.g., inhibiting dopamine synthesis; (Perez et al., 2002)) – making it a candidate for addiction research. In regards to alcohol, Foroud et al. (Foroud et al., 2007), identified haplotypes of SNPs in SNCA that were associated with alcohol craving, but not DSM-IV alcohol dependence diagnosis, supporting our argument here that the study of individual symptoms for AUD may at times point sources of liability that may be overlooked when studying only the shared variance across AUD symptoms. More recently, our analysis of genes in the dopamine pathway (i.e., DRD1, DRD2, DRD3, DRD4, SLC6A3, as well as SNCA) also suggested common and specific effects from variants in these genes across craving and alcohol dependence (i.e., without craving) (Agrawal et al., 2013). It is important to note that for the current analyses, craving was assessed using a single item (strong desire to use so couldn’t think of anything else). Prior work has contrasted the contributions of varying definitions of craving on AUD diagnosis (Keyes et al., 2011); the NESARC includes two items that effectively separate “strong desire or urge” from “couldn’t think about anything else” with the former being more commonly reported in population samples than the latter. However, “strong desire” is incorporated into the

<table>
<thead>
<tr>
<th>Symptom</th>
<th>Factor Loading [95% CI]</th>
<th>% Total Genetic Variance Explained</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sx1: Recurrent use resulting in failure to fulfill major roles</td>
<td>0.93 [0.92, 0.94]</td>
<td>87%</td>
</tr>
<tr>
<td>Sx2: Recurrent use in situations in which it is physically hazardous</td>
<td>0.81 [0.80, 0.83]</td>
<td>66%</td>
</tr>
<tr>
<td>Sx3: Continued use despite persistent or recurrent social problems</td>
<td>0.86 [0.85, 0.87]</td>
<td>74%</td>
</tr>
<tr>
<td>Sx4: Tolerance</td>
<td>0.74 [0.73, 0.76]</td>
<td>53%</td>
</tr>
<tr>
<td>Sx5: Withdrawal</td>
<td>0.80 [0.79, 0.82]</td>
<td>64%</td>
</tr>
<tr>
<td>Sx6: Taken in larger amounts or over a longer period than was intended</td>
<td>0.83 [0.82, 0.84]</td>
<td>69%</td>
</tr>
<tr>
<td>Sx7: Persistent desire to cut down or control alcohol use</td>
<td>0.90 [0.89, 0.91]</td>
<td>81%</td>
</tr>
<tr>
<td>Sx8: A great deal of time spent to obtain/use/recover from alcohol</td>
<td>0.76 [0.75, 0.78]</td>
<td>58%</td>
</tr>
<tr>
<td>Sx9: Given up or cut back on important activities in order to drink</td>
<td>0.85 [0.84, 0.86]</td>
<td>72%</td>
</tr>
<tr>
<td>Sx10: Continued to use alcohol despite knowledge physical or psychological problems</td>
<td>0.91 [0.90, 0.92]</td>
<td>83%</td>
</tr>
<tr>
<td>Sx11: Craving</td>
<td>0.61 [0.59, 0.64]</td>
<td>37%</td>
</tr>
</tbody>
</table>

Table showing standardized factor loadings of the exploratory genetic factor analysis along with 95% confidence intervals and squared standardized factor loading (i.e., percent of genetic variance explained).
parsimonious model. Altogether, when considering these factors, the pattern of results provide preliminary evidence to suggest that studying the shared liability across all of the DSM symptoms is a more genetically sensitive (i.e., evidencing a moderate heritability (0.30-0.60)) and parsimonious phenotype, since the loci likely reflects the lowest common denominator/factors for AUD.

In conclusion, we discovered that the APA's DSM-5 definition of alcohol-related problems is a heritable phenotype with varying genetic effects across the individual symptoms with both shared and non-shared genetic variance between them. Though tentative and in need of replication in larger samples, these findings lend support to the use of composite scores, such as factor scores or symptom count as phenotype, as well as the application of genomic structural equation model methods in future studies.

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Contributors

Authors Palmer, Agrawal, and Keller contributed to the overall design of the study and analytical plan. Analyses were carried out by Palmer, Chou, and Brick. Palmer wrote the first draft of the manuscript which was systematically reviewed and edited by Agrawal, Knopik, McGearry, Bierut, Hartz, Schuckit, and Johnson. All authors contributed to and approved of the final version of the manuscript.

Declaration of Competing Interest

No conflict declared.

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Appendix A. Supplementary data

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