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Title: Polygenic contributions to alcohol use and alcohol use disorders across population-based and clinically ascertained samples

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ABSTRACT

BACKGROUND

Studies suggest that alcohol consumption and alcohol use disorders have distinct genetic backgrounds.

METHODS

We examined whether polygenic risk scores (PRS) for consumption and problem subscales of the Alcohol Use Disorders Identification Test (AUDIT-C, AUDIT-P) in the UK Biobank (UKB; N=121,630) correlate with alcohol outcomes in four independent samples: an ascertained cohort, the Collaborative Study on the Genetics of Alcoholism (COGA; N=6,850), and population-based cohorts: Avon Longitudinal Study of Parents and Children (ALSPAC; N=5,911), Generation Scotland (GS; N=17,461), and an independent subset of UKB (N=245,947). Regression models and survival analyses tested whether the PRS were associated with the alcohol-related outcomes.

RESULTS

In COGA, AUDIT-P PRS was associated with alcohol dependence, AUD symptom count, maximum drinks ($R^2=0.47\% - 0.68\%$, $p=2.0e-8 - 1.0e-10$), and increased likelihood of onset of alcohol dependence (Hazard Ratio =1.15, $p=4.7e-08$); AUDIT-C PRS was not an independent predictor of any phenotype. In ALSPAC, the AUDIT-C

PRS was associated with alcohol dependence ($R^2=0.96\%$, $p=4.8e-6$). In GS, AUDIT-C PRS was a better predictor of weekly alcohol use ($R^2=0.27\%$, $p=5.5e-11$), while AUDIT-P PRS was more associated with problem drinking ($R^2=0.40\%$, $p=9.0e-7$). Lastly, AUDIT-P PRS was associated with ICD-based alcohol-related disorders in the UKB subset ($R^2=0.18\%$, $p<2.0e-16$).

CONCLUSIONS

AUDIT-P PRS was associated with a range of alcohol-related phenotypes across population-based and ascertained cohorts, while AUDIT-C PRS showed less utility in the ascertained cohort. We show that AUDIT-P is genetically correlated with both use and misuse and demonstrate the influence of ascertainment schemes on PRS analyses.

INTRODUCTION

Alcohol use, across the lifespan, can be characterized by a series of transitions: from early experimentation to hazardous drinking and possible development of alcohol use disorders (**AUD**), or to non-problematic alcohol intake. Some of these transitions are developmentally salient: for instance, hazardous drinking is common during adolescence and early adulthood, but does not always portend problematic use in later adulthood (Degenhardt et al., 2013; Wennberg, Andersson, & Bohman, 2000). Similarly, individuals with AUD might remit, even to abstinence, or persist into later life (McCutcheon et al., 2012; Trim, Schuckit, & Smith, 2013).

Furthermore, alcohol-related behavior can be broadly disarticulated into two components – the extent to which an individual consumes alcohol and the potential problems that they experience related to their intake. For instance, AUD are characterized not by excessive alcohol consumption but by measurable physiological changes that accompany addiction (e.g. withdrawal, tolerance) as well as loss of control over drinking and drinking despite physical and emotional impairment (American Psychiatric Association, 2013). These complexities underlying the life course of alcohol use and misuse pose challenges in the dissection of etiology.

Most stages of alcohol involvement include a heritable contribution (twin $h^2 = 40-70\%$), although the magnitude of these genetic effects varies considerably across development (Edwards et al., 2017; Enoch, 2006; Pagan et al., 2006). Common genetic variants from genome-wide association studies (**GWAS**) explain 4-13% of the phenotypic variance in alcohol use and misuse (Clarke et al., 2017; Kranzler et al., 2019; Liu et al., 2019; Sanchez-Roige et al., 2017; Schumann et al., 2016;

Walters et al., 2018). The largest GWAS of alcohol dependence to date (Kranzler et al., 2019) suggests that the genetic correlation between alcohol consumption (units per week) and ICD coded AUD is variable (r_g ranging from 0.54 (beer/cider) to 0.004 (champagne/white wine)).

A recent study (Sanchez-Roige et al., 2018) conducted GWAS of both the consumption and problem subscales of the Alcohol Use Disorders Identification Test (**AUDIT**), a short screener for drinking in the past year (Saunders et al., 1993). In the UK Biobank sample, genetic liability to the consumption subscale (**AUDIT-C**; three items with information pertaining to alcohol consumption) was positively correlated with educational achievement and unrelated to psychopathology whereas liability to the problem subscale (**AUDIT-P**; seven items with information pertaining to alcohol problems) was negatively correlated with educational achievement and positively correlated with psychopathology. These findings are consistent with the lack of genetic correlation between psychiatric illness and genetic liability to alcohol consumption in the largest GWAS of the trait (drinks per week; Liu et al., 2019). In contrast, two recent studies suggest moderate genetic correlations between AUD and consumption indices, including the AUDIT-C (e.g., $r_g = 0.52$, $p = 2.40e-42$) (Kranzler et al., 2019; Marees et al., 2019), while another study found that polygenic risk scores (**PRS**) for past week alcohol consumption predicted a modest but significant amount of variance in AUD (e.g., $R^2 = 0.56\%$; Johnson et al., 2019), suggesting that the genetic correspondence between recent consumption and dependence may be complicated by several factors, including the characteristics of the sample, and the nature of the assessment (e.g., alcohol quantity vs. frequency (Marees et al., 2019)).

In this study, we sought to examine the extent to which PRS derived from the AUDIT-C and AUDIT-P GWAS predicted variance in multiple aspects of alcohol use and misuse, ranging from levels of alcohol consumption, hazardous drinking, and AUD, in four independent samples that vary in their age and ascertainment scheme. PRS are scores that represent an individual's genetic liability for a certain trait or disorder, created by aggregating the effects of many risk variants for the phenotype of interest, weighting the effect sizes by the number of effect alleles an individual carries at each locus. Prior evidence indicates that association between PRS and phenotype in the target sample is improved when both the discovery and target samples for PRS analyses are derived using similar ascertainment strategies (Savage et al., 2018). Based on a study showing positive correlations between psychopathology and AUDIT-P (but not AUDIT-C; Sanchez-Roige et al., 2018), we hypothesized that AUDIT-P PRS would be more closely related to liability to AUD than would AUDIT-C PRS, which would be more closely related to aspects of alcohol consumption (e.g., regular consumption, units per week). We also hypothesized that associations with AUDIT-C would be stronger in the youngest sample while the AUDIT-P would be more predictive of drinking in older, ascertained samples in which problem drinking is more established. While there have been some recent studies examining the genetic overlap between alcohol consumption and indices of problem drinking (e.g., Johnson et al., 2019), none have yet compared the performance of consumption (AUDIT-C) versus problem drinking (AUDIT-P) PRS across multiple samples. Taken together, the current analyses demonstrate how genetic findings derived from a simple and fast screening tool could serve to outline the polygenic

underpinnings of different stages of alcohol use and problems in diversely ascertained samples.

MATERIALS AND METHODS

Discovery GWAS

The summary statistics of the recent GWAS of AUDIT-C and AUDIT-P in 121,604 unrelated individuals of European ancestry from the UK Biobank cohort (Sanchez-Roige et al., 2018) were used to construct PRS for alcohol-related outcomes in four independent target samples. The UK Biobank (**UKB**, www.ukbiobank.ac.uk) is a population-based sample of 502,629 participants who were recruited from 22 assessment centers across the United Kingdom from 2006 to 2010 (Bycroft et al., 2018); the AUDIT was delivered as part of the online Mental Health Questionnaire follow-up (Davis et al., 2018). Discovery GWAS analyses were performed using BGENIE (Bycroft et al., 2018), version 1.1, with AUDIT scores (AUDIT-C score, and AUDIT-P score, tested independently) as the outcome variable and age, sex, genotyping array, and the first 20 principal components derived from genotype data as covariates to account for any remaining population stratification. Further details regarding the discovery GWAS are provided in the **Supplemental Materials**.

Target sample demographics and characteristics

Collaborative Study on the Genetics of Alcoholism

The Collaborative Study on the Genetics of Alcoholism (**COGA**) (Begleiter et al., 1995; Nurnberger et al., 2004; Schuckit et al., 2018) includes probands meeting criteria for alcohol dependence, their family members, and community control

families. There were 7,645 participants of European ancestry with genotype data available as well as data on alcohol-related phenotypes; of these, 6,850 were ≥ 20 years (the mean age of onset of alcohol dependence) and reported lifetime alcohol use.

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Avon Longitudinal Study of Parents and Children

The Avon Longitudinal Study of Parents and Children (**ALSPAC**) recruited 14,541 pregnant women residing in Avon, UK, with expected dates of delivery April 1, 1991, to December 31, 1992. The enrolled sample consisted of 14,775 children. Additional details are provided in the **Supplemental Materials**. For the current analyses, phenotypic data were available for up to 5,911 participants, depending on the phenotype and wave of assessment. Ethical approval for the study was obtained from the ALSPAC Ethics and Law Committee and the Local Research Ethics Committees.

Generation Scotland

The Scottish Family Health Study (**GS**) is a family-based cohort recruited from the general population of Scotland from 2006-2011 (N=24,084) (Smith et al., 2013). A subset of GS participants responded to a re-contact in 2015 (Navrady et al., 2018) and provided additional information on mental health, which included information on alcohol misuse (N = 17,461).

UK Biobank

Individuals who were part of the AUDIT discovery GWAS (Sanchez-Roige et al., 2018) were removed from the UKB target sample, leaving 245,947 individuals with phenotypic data for these analyses.

Alcohol use measures

A summary of the alcohol-related outcomes included in the analyses is presented in **Table 1**. Across all samples, participants who did not endorse lifetime alcohol use were removed to account for those with high genetic liability who elect not to drink for personal or socio-cultural reasons, or were not old enough to have had a first drink. Each target sample used a different assessment. Details regarding each assessment are available in the **Supplementary Materials**. Briefly, COGA participants were administered the Semi-Structured Interview for the Genetics of Alcoholism (SSAGA; (Bucholz et al., 1994)), which includes diagnostic assessments for DSM-IV and DSM-5 AUD as well as other aspects of alcohol use. ALSPAC participants completed the AUDIT questionnaire, and DSM-IV alcohol dependence was derived for 4,328 participants, using symptom-level items adapted from (Kendler, et al., 1992). In GS, participants reported on their alcohol intake (units per week) using a pre-clinical questionnaire, and were also administered the CAGE (Cut-Annoyed-Guilty-Eye) screener (Ewing, 1984). In the UKB, ICD-9 and 10 codes for AUD were derived from linkage to hospital inpatient records. Phenotypic correlations between alcohol use measures across the samples are shown in **Supplementary Table 16**.

Genotypes

Details on genotyping and quality control are shown in the **Supplementary Materials**.

Polygenic risk score construction

For all samples, PRS were coded for every individual by multiplying an individual's number of effect alleles at a particular SNP by that SNP's effect size (beta) from the discovery GWAS (Sanchez-Roige et al., 2018) then averaging across SNPs to create one score per person. The discovery GWAS summary statistics were clumped using PLINK (Chang et al., 2015; Purcell et al., 2007) using the linkage disequilibrium (**LD**) pattern from the 1000 Genomes European (Consortium, 2015) reference sample, with an LD threshold of $r^2 \geq 0.25$ and a 500 kb physical distance. PRS were constructed for 8 thresholds ($p_T < 0.0001$, $p_T < 0.001$, $p_T < 0.01$, $p_T < 0.10$, $p_T < 0.20$, $p_T < 0.30$, $p_T < 0.40$, $p_T < 0.50$) and standardized using the scale function in R within each sample to ease interpretation of effect sizes. For ALSPAC and COGA, PRS were generated using PLINK; for UKB and GS samples, PRS were created using PLINK implemented in PRS-ice2 (Euesden et al., 2014) using the same R^2 thresholds for clumping and the same p-value thresholds.

Statistical Analyses

Linear and logistic regression models were used to examine whether AUDIT-C and AUDIT-P PRS were associated with each of the alcohol-related phenotypes (see **Table 1**), and to determine which PRS threshold (i.e., p_T) was most predictive of each measure based on the p-value and observed (linear) or pseudo (logistic) R^2 . For the cohorts of unrelated individuals (ALSPAC and UKB), the partial R^2 was extracted from linear regression models for continuous traits, while Nagelkerke's

pseudo-R² was extracted from logistic regression models for binary traits. For the cohorts that employed mixed-effect models to account for within-sample relatedness (COGA and GS), variance explained by the PRS in the continuous outcomes (e.g. MaxDrinks, CAGE) was calculated by multiplying the PRS by its regression coefficient and dividing the variance of that value by the variance of the outcome to derive a coefficient of determination between 0 and 1 (Nakagawa & Schielzeth, 2012); in COGA, the ‘MuMIn’ package in R was used to calculate marginal R² for the logistic mixed-effect models for the binary outcomes (Barton, 2011). Once the most predictive AUDIT-C and AUDIT-P PRS thresholds were determined in the single-PRS models (n.b. p_T was allowed to vary between AUDIT-C and AUDIT-P PRS), the most predictive PRS for both AUDIT-C and AUDIT-P PRS were simultaneously entered into a joint regression model along with covariates (results for single-PRS models are available in **Supplemental Table 1-12**). We primarily report on the results of the joint (AUDIT-C PRS + AUDIT-P PRS) regression models, as we wished to examine the relative contribution of each AUDIT subscale PRS while controlling for the other subscale PRS. As the primary analyses consisted of 2 tests (AUDIT-C PRS and AUDIT-P PRS) for each of the 12 outcomes, and the number of independent tests across the PRS p-value thresholds was estimated to be approximately 5 (calculated using spectral decomposition, via the matSpD.R R script (Nyholt, 2004)), we corrected for 120 tests using a Bonferroni p -value = 0.0004.

COGA: Sex, 4 ancestral principal components (chosen via visual inspection of a scree plot of the eigenvalues), total number of interviews, birth cohort (born 1890-1929, 1930-1949, 1950-1969, \geq 1970), and array type (see **Supplemental**

Materials) were included as fixed effects, while the family identifier and recruitment site were included as random effects (family nested within site).

ALSPAC: In keeping with standard practice for genetic analyses in ALSPAC, sex and 10 ancestral principal components were included as covariates. ALSPAC assessments are age-specific, therefore age was not included as a covariate.

GS: Analyses were performed in AS-REML-R fitting pedigree information as a random effect to control for relatedness in the sample. Fixed effects included age, sex and 4 multidimensional scaling components to correct for population stratification. _

UKB: Age, sex and 10 ancestry principal components were used as covariates.

Cox proportional hazards survival models for onset of hazardous drinking (AUDIT total score ≥ 8) in ALSPAC and DSM-IV alcohol dependence in COGA were fit to the data with identical covariates as the regression analyses. Those who did not meet criteria for hazardous drinking or alcohol dependence were censored at their age at last interview (since age at birth). For ease of comparison across the two samples, the same PRS threshold ($p_T < 0.5$) was used in both COGA and ALSPAC and for both the AUDIT-C and AUDIT-P PRS. Violations of the proportional hazards assumption for the PRS were tested using scaled Schoenfeld residuals. All analyses were conducted in R (R Core Team, 2017). In COGA, family identifier and recruitment site were included in the survival models as a nested frailty term (i.e. `(1 | site/family)`), using the “coxme” package for mixed-effect Cox proportional hazards models

containing Gaussian random effects (Therneau, 2018). ALSPAC data were analyzed using the “survival” and “survminer” packages in R (Kassambara et al., 2017; Therneau & Lumley, 2015)

For illustrative purposes, PRS were also split into quartiles for COGA, ALSPAC, and UKB, and the odds of having an AUD diagnosis (DSM-IV alcohol dependence in COGA and ALSPAC, ICD-9/ICD-10 alcohol related codes in UKB) was calculated in each quartile using the lowest quartile as the reference group.

RESULTS

Target sample demographics and characteristics

The prevalence of each alcohol use measure for each target sample is shown in **Table 1**. The SNP-heritability of AUDIT-C was 11%, while the SNP-heritability of the AUDIT-P score was 9%, and the genetic correlation (i.e., $SNP-r_g$) between them was $r_g = 0.70$ (as reported in the original paper (Sanchez-Roige *et al.*, 2018)). See **Supplemental Table 15** for the number of SNPs in each PRS threshold.

COGA: 33.8% of the participants (N = 2,318) met DSM-IV criteria for alcohol dependence during their lifetime, with 77.6 % of those endorsing ≥ 6 (i.e., severe) DSM-5 AUD criteria. A median of 14.0 drinks (SD = 12.9; ~96 grams of alcohol, or 14 standard US drinks) in a single 24-hour period was reported (i.e., MaxDrinks). 90.0% of the participants reported drinking at least once a month for 6 consecutive months or longer (i.e., monthly alcohol use).

ALSPAC: By age 23, 11.2% (N = 484) had met DSM-IV criteria for alcohol dependence within the past year at any of the three assessment waves, while

66.2% reported hazardous drinking (AUDIT total score ≥ 8). The median MaxDrinks was 15 UK units (SD = 12.4; approximately 120 grams of alcohol, or 8.5 standard US drinks).

GS: Mean alcohol intake was 10.9 units per week (SD = 12.8; approximately 87.2 grams of alcohol, or 6.2 standard US drinks; N = 17,461) and the mean CAGE score was 0.6 (SD = 0.93).

UKB: There were 4,141 (1.68%) individuals in the subset of UKB selected for analysis who had at least one ICD-9 or ICD-10 code assigned as a primary or secondary diagnosis of alcohol related disorders during time as a hospital inpatient.

Associations between AUDIT PRS and consumption measures

COGA: As shown in **Table 2**, only the AUDIT-P PRS was significantly associated with monthly alcohol use ($R^2 = 0.65\%$, $p = 1.24e-4$) and MaxDrinks ($R^2 = 0.47\%$, $p = 2.00e-8$). We also converted the pseudo- R^2 values for all binary outcomes to R^2 on the liability scale (Lee *et al.*, 2012), but as estimates did not differ, these results are not presented.

ALSPAC: AUDIT-C and AUDIT-P PRS accounted for increasing proportions of phenotypic variance in AUDIT-C and AUDIT-P scores as the sample aged, from $\sim 0.50\%$ at age 16 to 1.12% at age 23 (see **Supplementary Tables 8-9** for complete results). Here we focus on results for age 23 for comparability to other samples, which consisted predominantly of adults. No association survived the correction for multiple tests. The strongest associations observed were for AUDIT-C and AUDIT-P PRS, which were both associated with higher AUDIT-C scores in ALSPAC at age 23, each explaining 0.60% of the variance ($p = 1.06e-3$ and $5.9e-4$,

respectively), and between AUDIT-P PRS and MaxDrinks ($R^2 = 3.30\%$, $p = 1.59e-3$); a 1-SD increase in AUDIT-P PRS corresponds to a predicted increase of 0.54 drinks. GS: The AUDIT-P PRS explained 0.15% of the variance in units per week ($p = 4.7e-7$) and 0.40% of the variance in CAGE scores ($p = 9.0e-7$), while the AUDIT-C PRS explained 0.27% of the variance in units per week ($p = 5.5e-11$).

Associations between AUDIT PRS and alcohol problem measures

COGA: As shown in **Table 2**, only the AUDIT-P PRS was significantly associated with DSM-IV alcohol dependence ($R^2 = 0.68\%$, $p = 4.55e-9$) and DSM-5 AUD symptom count ($R^2 = 0.67\%$, $p = 1.01e-10$). For individuals in the top quartile of AUDIT-P PRS risk, the odds of being diagnosed with alcohol dependence was 1.74 (95% confidence intervals (C.I.) = 1.47-2.07) compared to those in the lowest quartile, while being in the top AUDIT-C PRS quartile was associated with an odds ratio (OR) of 1.30 (95% C.I. = 1.10 -1.54) compared to those in the lowest quartile (**Figure 1**). Comparison of the OR and their 95% C.I. suggest that for AUDIT-C, those in the 4th quartile are at significantly elevated likelihood of AUD compared to those in lowest quartile, though there were overlapping 95% C.I. between the 2nd, 3rd, and 4th quartiles. In contrast, for AUDIT-P, likelihood of AUD escalates at the mid-point, with those in the both the 3rd and 4th quartiles being at significantly greater risk of AUD than those in the lowest quartile of AUDIT-P PRS.

ALSPAC: Only AUDIT-P PRS were associated with AUDIT-P scores, accounting for 1.1% of the variance ($p = 9.05e-6$). AUDIT-C PRS and AUDIT-P PRS were both positively associated with DSM-IV alcohol dependence, explaining 1.0% ($p = 4.81e-6$) and 0.50% ($p = 5.75e-4$) of the variance, respectively, although only the AUDIT-C PRS passed multiple testing corrections. Both PRS scores were also positively

associated with DSM-IV alcohol dependence symptom count ($R^2 = 0.8\%$ for both; AUDIT-C $p = 3.59e-5$, AUDIT-P $p = 3.53e-5$). For individuals in the top quartile of AUDIT-C PRS, the odds of being diagnosed with AUD was 1.73 (95% C.I. = 1.29-2.33) compared to those in the lowest quartile, while being in the top AUDIT-P PRS quartile was associated with an OR of 1.25 (95% C.I. = 0.93 -1.68) compared to those in the lowest quartile (**Figure 1**). Due to wider 95% C.I. (suggesting imprecision potentially due to smaller sample size and/or less of a dose-response association pattern), it was difficult to draw comparisons of the ORs across quartiles in the ALSPAC sample. For AUDIT-C, differentiation in likelihood of AUD appeared to occur at the 3rd quartile, while for AUDIT-P, there was no clear distinction in the OR across the 2nd, 3rd and 4th quartiles.

UKB: Positive associations with AUD-related ICD codes were observed for both AUDIT-P ($R^2 = 0.18\%$, $p < 2e-16$) and AUDIT-C ($R^2 = 0.04\%$, $p = 8.9e-5$) PRS. For individuals in the top quartile of AUDIT-P PRS, the odds of having a mental or behavioral disorder due to alcohol was 1.45 (95% C.I. = 1.37-1.54) compared to those in the lowest quartile. For AUDIT-C PRS, being in the top PRS quartile was associated with an odds-ratio of 1.11 (95% C.I. = 1.02-1.20) compared to those in the lowest quartile (**Figure 1**). In the UKB, likelihood of AUD across quartiles seemed to follow a dose-response pattern with increasing OR for each quartile of polygenic risk for both AUDIT-C and AUDIT-P (though with overlapping C.I. for AUDIT-C), with the 4th quartile of AUDIT-P PRS being statistically significantly more likely to be diagnosed with AUD relative to the 3rd quartile.

Associations between AUDIT PRS and onset of hazardous drinking and alcohol dependence

COGA: The AUDIT-P, but not the AUDIT-C, PRS was associated with increased hazards of alcohol dependence onset (AUDIT-P HR = 1.15, 95% CI = 1.10-1.20, $p = 4.7e-08$; **Supplemental Table 13**). Those in the top quartile of AUDIT-P PRS were at considerably higher risk for onset of alcohol dependence (HR = 1.50, 95% CI = 1.36-1.64, $p = 9.7e-10$) relative to any other group; the risk attributable to the top AUDIT-C PRS was lower in magnitude (HR = 1.27, 95% CI = 1.14-1.40, $p = 1.8e-4$). ALSPAC: Only the AUDIT-C PRS was associated with onset of hazardous drinking, although not significant after multiple testing corrections (**Supplemental Table 14**; AUDIT-C HR = 1.06, 95% CI = 1.02-1.10, $p = 2.28e-3$).

DISCUSSION

The course of alcohol use and misuse consists of several stages, each with polygenic and environmental underpinnings. At earlier ages, how much and how often a person drinks often serves as an indicator of their genetic susceptibility to alcohol misuse (Dick et al., 2011; Irons et al., 2015). As an individual progresses towards maladaptive alcohol intake, a unique genetic susceptibility to physiological and psychological impairments related to alcohol misuse may unfold (Kendler et al., 2011). Our goal was to outline the extent to which PRS derived from a discovery GWAS of the AUDIT consumption (AUDIT-C) and problem (AUDIT-P) subscales predicted variance in a range of alcohol use behaviors across samples that were variously ascertained. With few exceptions, polygenic liability to AUDIT-P was a superior predictor of indices of heavy drinking and alcohol problems compared to the AUDIT-C.

Overall, across several ascertainment strategies, AUDIT-P PRS outperformed AUDIT-C as a predictor of both normative and disordered drinking (see **Figure 2** for an overview). For example, AUDIT-P PRS predicted up to 3.3% of the variance in the lifetime maximum number of drinks consumed in a 24-hour period (although this was a clear outlier in terms of variance explained), 1.1% of the variance in AUDIT-P scores, and 0.7% of the variance in DSM-5 AUD symptom counts. Exceptions to the superior performance of AUDIT-P PRS were observed within ALSPAC, where AUDIT-C PRS explained higher proportions of variance than AUDIT-P PRS in some adolescent AUDIT-C and AUDIT-P assessments (prior to age 23; see **Supplemental Materials**). AUDIT-C PRS also accounted for more variance in DSM-IV AD in ALSPAC.

Even indices of alcohol consumption, such as lifetime maximum drinks in 24 hours and monthly alcohol use, were more closely related to the genetic liability indexed by AUDIT-P PRS than AUDIT-C PRS. However, this difference was most notable in COGA, for which drinking for at least 6 months during the lifetime represents a highly heterogeneous group of individuals, which may include both non-problem drinkers and those with AUD. As expected, and in line with prior studies (Mies et al., 2018; Sanchez-Roige et al., 2018), AUDIT-P outperformed AUDIT-C for measures of problem drinking within each sample (i.e., COGA, ALSPAC, GS and UKB; see **Table 2**), confirming that the AUDIT-P is more closely related to genetic susceptibility to alcohol misuse than AUDIT-C.

Overall, the variance explained by PRS was low, even when the discovery and target phenotypes were identical (e.g. AUDIT scores in ALSPAC). These estimates are nonetheless consistent with other PRS studies (Mies et al., 2018; Savage et al.,

2018), and the proportion of explained variance may improve as the discovery GWAS get larger. In agreement with previous studies (Savage et al., 2018), we demonstrated that ascertainment is likely to be a crucial factor in polygenic prediction analyses. For instance, while the AUDIT-P PRS explained a larger amount of variance in dependence in COGA than in ALSPAC, the AUDIT-C PRS was not a significant predictor for any phenotype in COGA over and above the effect of AUDIT-P PRS, suggesting that the effect sizes for AUDIT-C from a volunteer cohort (UKB) may not be optimal for capturing genetic risk to AUD in high-risk families, such as those in COGA. AUDIT-C associations in COGA using other addiction-enriched samples, such as the Million Veteran Program (Kranzler et al., 2019), where the genetic correlation between AUDIT-C and AUD is high, will likely provide insights into the extent of the effect of ascertainment differences in discovery and target samples. Interestingly, variance explained by either PRS was the highest in ALSPAC, a population-based cohort that is noticeably younger than the discovery sample (age 39-79 years). Lastly, the variance explained for AUD in the (independent) subset of UKB participants itself was markedly lower than in ALSPAC. Here, it is worth recognizing that AUDIT is a past-year screener for alcohol consumption/problems, and there may have been individuals in the original AUDIT GWAS with low scores who were formerly problem drinkers. Furthermore, using ICD codes derived from hospital records as a proxy for AUD in the UKB may have resulted in false negatives; some of the 'controls' could have been problem drinkers but had not been diagnosed with an alcohol related condition as a hospital inpatient. Thus, the lower prediction in the UKB sample may reflect the instruments used to measure AUD in both the discovery and target GWAS.

Our findings should be viewed in the context of several limitations. First, the AUDIT is a past-year screener for alcohol consumption and problems, and thus may not capture individuals who had alcohol use problems in the past. Second, the AUDIT discovery GWAS was performed in a subset of the UKB that consisted of individuals who responded to an email request to complete a mental health follow-up survey. A recent study of the genomic profiles of these individuals has found them to represent lower polygenic risk for psychopathology and greater polygenic load for educational achievement, suggesting selection bias in the discovery sample itself (Adams et al., 2018). In addition, these individuals had relatively low endorsement of AUDIT-P items, which might have limited the statistical power of the discovery GWAS to capture the full spectrum of problem drinking and, in turn, limited our ability to compare across the AUDIT-C and AUDIT-P. Third, as the discovery GWAS was conducted on individuals of primarily European descent, these PRS could not be applied to other ancestral groups, reflecting a research gap in the dearth of large discovery GWAS of non-European populations (Martin et al., 2017). Results from the recent AUDIT-C GWAS by Kranzler et al (2019), which includes 57,340 African-Americans, for instance, will be instrumental in bridging this gap. Fourth, it is likely that the pattern of results reflects a complex interplay between developmental effects, ascertainment strategy, the psychometrics of the assessment instruments, and even population selection effects. For instance, distinctions between ALSPAC and COGA could be attributed to the higher median age of COGA participants, or even drinking patterns prevalent in earlier birth cohorts that are represented within COGA. Potential moderation of PRS by sex are also possible; however, power for such exploratory analyses is limited and beyond the scope of this study. Fifth, although this study examined a range of alcohol use phenotypes (from a measure of

monthly alcohol use to dependence), the cross-sectional design cannot capture potential longitudinal changes in PRS association across the progression of stages of alcohol use. A sixth limitation is that we did not control for any measures of socioeconomic status (SES) in our models, although SES has been shown to be a potential moderator of polygenic risk for alcohol consumption (Clarke et al., 2016; Barr et al., 2018), as the measures available varied widely and inconsistently across our target samples. The imbalance of cases and controls in the UK Biobank target sample is another limitation; this imbalance would be more likely to seriously bias a discovery GWAS, but may still have confounded the regression analyses in our target sample. A final limitation is that the strength of PRS associations varied across p-value thresholds (p_T ; see **Supplementary Tables 1-12**); because we chose the best-performing p_T for each model, as is typical for the pruning+thresholding (P+T) method of polygenic risk score analyses (Euesden et al., 2014), we run the risk of over-fitting.

Collectively, these analyses demonstrate that much of the polygenic liability to alcohol use and misuse remains unaccounted for – we anticipate that much larger GWAS of phenotypes representing disordered drinking, in conjunction with other risk indices (e.g., PRS for negative affect, a hallmark of later stages of AUD) will be necessary to explain additional variance. Importantly, as the use of PRS becomes increasingly common in attempts to dissect the transitions from experimentation, to regular or problem use and further, to AUD, differences between the discovery sample and target sample(s) in age, ascertainment, and other characteristics will need to be considered in the interpretation of findings. In addition, our findings highlight the high degree of heterogeneity and polygenicity underlying alcohol use

and misuse – not only is there no “gene for” alcoholism, there is currently no robust “polygenic indicator for” clinical prediction of liability to problem drinking. Furthermore, while we focused on contrasting the genetic relationship of each individual AUDIT subscale with the variety of alcohol-related measures across our different samples, it should be noted that the proportion of variance explained in the alcohol-related outcomes is larger when we consider both the AUDIT-C and AUDIT-P PRS together, suggesting that multiple polygenic scores provide more utility than PRS for the individual sub-scales. The two scores are only modestly correlated and likely, some of this commonality is due to variants in *ADH1B* that exert a relatively large effect. Overall, our analyses suggest that even though it is far too small to be statistically significant, there is some incremental contribution of variance from inclusion of the AUDIT-C. Despite these caveats, our analyses demonstrate the feasibility of using a fast and simple screening questionnaire, the AUDIT, that assesses both consumption and problem drinking, to understand the course of alcohol use and misuse.

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TABLES

Table 1. Alcohol-related measures in each target sample. C, Continuous (linear regression model used); B, Binary (logistic regression model used)

Drinking measure		Definitions			
Trait	Continuou s/ Binary	COGA Ascertained for AUD	ALSPAC Population-based	GS Populatio n-based	UKB Population -based
Units per week (Mean, SD)	C	n/a	n/a	10.9 (12.8)	n/a
Alcohol use (N, %)	B	Ever drank once a month for 6 consecutive months or longer 6,163 (90.0%)	n/a	n/a	n/a
Maxdrinks (Median, SD)	C	Lifetime maximum drinks consumed in a single 24-hour period 14.0 (12.87)	Lifetime maximum drinks consumed in a single 24-hour period 15.0 (12.35)	n/a	n/a
Hazardous drinking (N, %)	B	n/a	AUDIT total score cutoff of 8+ at any age 3264 (66.2%)	n/a	n/a
AUDIT-C score (Mean, SD)	C	n/a	Sum of AUDIT items 1-3 Age 16 4.80 (2.59) Age 18 5.04 (2.26) Age 19 6.01 (2.50)	n/a	n/a

			Age 21 6.11 (2.43) Age 23 5.37 (2.31)		
AUDIT-P score (Mean, SD)	C	n/a	Sum of AUDIT items 4-10 Age 16 2.37 (3.18) Age 18 2.48 (2.97) Age 19 2.38 (2.89) Age 21 3.16 (3.66) Age 23 2.30 (2.93)	n/a	n/a
DSM alcohol dependence symptom count (Mean, SD)	C	Sum of 11 lifetime criteria for diagnosis of DSM-5 AUD 3.57 (3.68)	Sum of 7 criteria for diagnosis of DSM-IV AUD; maximum observed across waves Age 18 0.35 (0.91) Age 21 0.72 (1.30) Age 23 0.43 (1.01)	n/a	n/a
DSM-IV alcohol dependence diagnosis (N, %)	B	Clustering of ≥ 3 of 7 DSM-IV criteria within 12 months 2,318 (33.8%)	Clustering of ≥ 3 of 7 DSM-IV criteria within 12 months at any wave of assessment (age 18, 21, or 23) 484 (11.2%)	n/a	n/a
ICD-9 and ICD- 10 alcohol dependence diagnosis (N, %)	B	n/a	n/a	n/a	ICD-9 code of 303* or ICD-10 code of F10* 4,141 (1.68%)
CAGE scores (Range 0-4; Mean, SD)	C	n/a	n/a	0.603 (0.93)	n/a

Table 2. Associations between AUDIT-C and AUDIT-P PRS and multiple indices of alcohol use in COGA, UKB, ALSPAC and GS samples. The PRS threshold (p_T) that most strongly predicted each outcome (based on highest R^2) is presented. Results significant at $p < 0.0004$ are bolded. Complete results are available in Supplemental Tables 1-12.

Outcome	Sample	AUDIT-C PRS				AUDIT-P PRS			
		p_T	R^2	Beta (SE)	p-value	p_T	R^2	Beta (SE)	p-value
Units per Week	GS	<0.1	0.27 %	0.052 (0.008)	5.50E-11	<0.01	0.15 %	0.039 (0.008)	4.70E-07
Alcohol use	COGA	<i><0.0001</i>	0.32 %	0.129 (0.046)	5.19E-03	<0.04	0.65 %	0.188 (0.049)	1.24E-04
AUDIT-C	ALSPAC (age 23)	<i><0.1</i>	0.59 %	0.066 (0.020)	1.06E-03	<i><0.01</i>	0.65 %	0.067 (0.020)	5.90E-04
Max Drinks	ALSPAC	<i><0.5</i>	1.54 %	0.389 (0.173)	2.44E-02	<i><0.2</i>	3.26 %	0.540 (0.171)	1.59E-03
	COGA	<i><0.001</i>	0.04 %	0.266 (0.147)	7.03E-02	<0.2	0.47 %	0.879 (0.156)	2.00E-08
CAGE	GS	<i><0.4</i>	0.19 %	0.044 (0.010)	8.60E-04	<0.2	0.40 %	0.063 (0.010)	9.00E-07
AUDIT-P	ALSPAC (age 23)	<i><0.2</i>	0.28 %	0.045 (0.020)	2.61E-02	<0.01	1.12 %	0.088 (0.020)	9.05E-06
Alcohol Dependence Symptom Count	ALSPAC (DSM-IV)	<0.3	0.84 %	0.091 (0.022)	3.59E-05	<0.01	0.84 %	0.086 (0.021)	3.53E-05
	COGA (DSM-5)	<i><0.1</i>	0.08 %	0.103 (0.045)	2.30E-02	<0.4	0.67 %	0.301 (0.046)	1.01E-10
Alcohol Dependence	ALSPAC (DSM-IV)	<0.3	0.96 %	0.240 (0.052)	4.81E-06	<i><0.01</i>	0.54 %	0.170 (0.050)	5.75E-04
	COGA (DSM-IV)	<i><0.0001</i>	0.14 %	0.078 (0.032)	1.44E-02	<0.4	0.68 %	0.199 (0.034)	4.55E-09
	UKB (ICD)	<0.0001	0.04 %	0.062 (0.016)	8.90E-05	<0.5	0.18 %	0.147 (0.020)	<2.00E-16

FIGURE CAPTIONS

Figure 1. Odds Ratios (OR) and 95% confidence interval for alcohol

dependence diagnosis by PRS quartiles in ALSPAC, COGA, and UKB. PRS

were split into quartiles and odds ratios calculated for case status for each quartile

of risk compared to quartile 1 (lowest). **A.** Alcohol dependence in ALSPAC was

coded by DSM-IV diagnosis (484 cases, 3,837 controls). **B.** Alcohol dependence was

coded via DSM-IV diagnosis in the COGA sample (2,318 cases, 4,532 controls). **C.** A

sample of European ancestry, unrelated British individuals who had ever drunk, had

ICD9 and ICD-10 codes available, and were not included in the discovery GWAS

were used (4,141 cases, 241,806 controls).

Figure 2. Overview of the magnitude of predicted variance (R^2 , %) across the alcohol-related phenotypes probed in the four independent samples (Generation Scotland, GS; ALSPAC at age 23, A23; Collaborative Study on the Genetics of Alcoholism, COGA; and UK Biobank, UKB). The color of the dots denotes AUDIT-C (light gray) or AUDIT-P (dark gray) PRS. Only the significant ($p < 0.0004$) associations are shown. C, AUDIT-C; P, AUDIT-P.