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# A genome-wide gene-by-trauma interaction study of alcohol misuse in two independent cohorts identifies PRKG1 as a risk locus

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## A genome-wide gene-by-trauma interaction study of alcohol misuse in two independent cohorts identifies *PRKG1* as a risk locus

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### Abstract

Traumatic life experiences are associated with alcohol use problems, an association that is likely to be moderated by genetic predisposition. To understand these interactions, we conducted a gene-

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#### Conflict of Interest

Dr. Stein has in the last three years been a consultant for Actelion Pharmaceuticals, Healthcare Management Technologies, Janssen, Pfizer, Resilience Therapeutics, Tonix Pharmaceuticals, and Oxeia Biopharmaceuticals. Dr. Kaufman has provided consultation to Pfizer and Merck Pharmaceutical Company to train investigators to assess bipolar disorder in youth. Dr. Kranzler has been an advisory board member, consultant, or CME speaker for Indivior, Lundbeck, and Otsuka. He is also a member of the American Society of Clinical Psychopharmacology's Alcohol Clinical Trials Initiative, which is supported by AbbVie, Alkermes, Ethypharm, Indivior, Lilly, Lundbeck, Pfizer, and XenPort. The other authors reported no biomedical financial interests or potential conflicts of interest.

by-environment genome-wide interaction study (GEWIS) of alcohol use problems in two independent samples, the Army STARRS (ASTARRS, N=16,361) and the Yale-Penn (N=8,084) cohorts. Because the two cohorts were assessed using different instruments, we derived separate dimensional alcohol misuse scales and applied a proxy-phenotype study design. In African-American subjects, we identified an interaction of *PRKGI* rs1729578 with trauma exposure in the ASTARRS cohort and replicated its interaction with trauma exposure in the Yale-Penn cohort (discovery-replication meta-analysis:  $z=5.64$ ,  $p=1.69 \times 10^{-8}$ ). *PRKGI* encodes cGMP-dependent protein kinase 1, which is involved in learning, memory, and circadian rhythm regulation. Considering the loci identified in stage-1 that showed same effect directions in stage-2, the gene ontology (GO) enrichment analysis showed several significant results, including calcium-activated potassium channels (GO:0016286;  $p=2.30 \times 10^{-5}$ ), cognition (GO:0050890;  $p=1.90 \times 10^{-6}$ ), locomotion (GO:0040011;  $p=6.70 \times 10^{-5}$ ), and Stat3 protein regulation (GO:0042517;  $p=6.4 \times 10^{-5}$ ). To our knowledge, this is the largest GEWIS performed in psychiatric genetics, and the first GEWIS examining risk for alcohol misuse. Our results add to a growing body of literature highlighting the dynamic impact of experience on individual genetic risk.

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## Introduction

Exposure to traumatic life events is associated with a variety of health risk behaviors, including alcohol use disorders (AUD).<sup>1, 2</sup> The heritability of AUD is approximately 50%.<sup>3</sup> Genome-wide association studies (GWAS) of AUD have identified several risk alleles.<sup>4</sup> Individuals with AUD likely present some complex trait risk mechanisms that are different from those of the general population.<sup>5, 6</sup> A recent phenome-wide analysis demonstrated that AUD risk alleles are associated with a wide range of physical and mental health consequences.<sup>7</sup> The environment also contributes to the predisposition to AUD, moderating the effects of risk alleles<sup>8</sup>.

Traumatic events affect genome regulation via different mechanisms;<sup>9, 10</sup> and it should be possible to identify specific genes that interact with traumatic experiences to moderate AUD risk. Previous studies focused on candidate stress-response genes, such as *5-HTTLPR*, *PER1*, and *FKBP5*,<sup>11-13</sup> and investigated how the exposure to life trauma interacts with risk alleles in relation to AUD. However, the candidate-gene approach has limited ability to identify the genetic basis of complex traits.<sup>14</sup> Conversely, GWAS of complex traits conducted in large cohorts have identified numerous risk alleles and some of the pathogenic mechanisms underlying genetically complex diseases. Similarly, genome-wide gene-by-environment interaction studies (GEWIS) can be useful in understanding how environmental factors interact with an individual's genetic background to regulate the predisposition to complex traits. However, to date, there have been few published GEWIS. One reason is that large cohorts that could be meta-analyzed are rarely ascertained using compatible criteria, and relevant differences can be present in the assessment of phenotypic outcome and environmental factors. Thus, it is difficult to investigate large cohorts evaluated with homogeneous assessments. Differences in phenotypic assessment can reduce the statistical power of meta-analysis and replication studies, especially for GEWIS, since they are performed using two kinds of phenotypic data with respect to both the outcome and an environmental factor.

Using data from a GWAS of AUD, we observed that risk alleles in *ADH1B*, one of the best-validated loci for alcohol drinking behaviors, show different associations with DSM-IV vs. DSM-5 AUD criteria.<sup>15</sup> To reduce error related to different phenotypic assessments, we performed a genome-wide gene-by-trauma interaction study considering the proxy-phenotype approach proposed by Rietveld and colleagues in 2014.<sup>16</sup> Specifically, we used the cohorts from the Army STARRS (ASTARRS) Initiative (N = 16,361) for stage-1 and the Yale-Penn sample (N = 8,084) for stage-2, which yielded a total sample of 24,445 individuals. To our knowledge this is the largest GEWIS performed in psychiatric genetics and the first examining risk for alcohol misuse.

## Subjects and Methods

### Army STARRS cohorts

Subjects investigated were selected from among the participants in the ASTARRS Initiative. All subjects gave written informed consent to participate. These procedures were approved by the Human Subjects Committees of all collaborating institutions.

**Sample**—Two study populations were included in the ASTARRS Initiative (Table 1). The New Soldier Study (NSS) includes soldiers at the start of their basic training at one of three Army installations. The Pre-Post Deployment Study (PPDS) is a multiple-wave panel survey that collected baseline data (time 0) from US Army soldiers in three brigade combat teams prior to their deployment to Afghanistan. Detailed information about the design and conduct of Army STARRS is available in a previous report.<sup>17</sup>

**Procedures**—Phenotypes for ASTARRS were obtained using a self-administered questionnaire, which included a computerized version of the Composite International Diagnostic Interview Screening Scales (CIDI-SC).<sup>18</sup> From the CIDI-SC assessment, we extracted information regarding lifetime trauma exposure and alcohol misuse.

**Lifetime Trauma Assessment**—Lifetime trauma exposure (i.e., exposed vs. unexposed) included reporting of any of the following experiences: serious physical assault; sexual assault or rape; witnessing someone being seriously injured or killed; discovering or handling a dead body; a life-threatening illness or injury; a disaster; any other experience that put the subject at risk of death or serious injury; murder of a close friend or relative; suicide of a close friend or relative; combat death of a close friend or relative; or accidental death of a close friend or relative. Further details on the trauma assessments were reported in our previous study.<sup>19</sup>

**Alcohol Use Assessment**—For the ASTARRS cohort, a dimensional measure of alcohol misuse was derived by summing responses to 13 items that assessed frequency and consequences of alcohol use including the array of alcohol misuse symptoms: 1) Frequency of Drinking; 2) Frequency of Binge Drinking; 3) Drinking Interfered with Responsibility; 4) Drinking Caused an Argument; 5) Drinking Resulted in Someone Getting Hurt; 6) Out of Control Drinking; 7) Arrested Due to Drinking; 8) Worried to Not Be Able to Drink; 9) Worried About Drinking; 10) Feel a Need to Cut Down; 11) Feel Annoyed by People Who Mention Drinking; 12) Feel Guilty About Drinking; 13) Drink First Thing in the Morning.

Respondents rated each symptom on a 5-point frequency scale that ranged from “never” through “every or nearly every day”. Supplemental Table 1 summarizes the items related to lifetime trauma exposure and alcohol-related symptoms. We included only subjects who reported having ever consumed an alcoholic drink (i.e., alcohol exposed). For PPDS subjects, we considered trauma and alcohol information reported at time 0 (within approximately six weeks prior to deployment).

**Genetics**—The NSS and PPDS samples (ASTARRS1, N=14,000) underwent genotyping using the Illumina OmniExpress and Exome array with additional custom content. An additional 2,361 NSS samples (ASTARRS2) were genotyped on the Illumina PsychChip array. Methods for genotyping, imputation, ancestry assignment, and principal component (PC) analysis were described previously.<sup>19, 20</sup>

### Yale-Penn Cohort

**Sample**—The subjects in the Yale-Penn cohort were recruited at five sites in the Eastern United States and were previously investigated in genetic studies of substance use disorders and other traits (Table 1).<sup>4, 21–24</sup> The institutional review board at each participating site approved the study and we obtained written informed consent from each participant.

**Trauma and Alcohol Use Assessment**—The Yale-Penn participants were evaluated using the Semi-Structured Assessment for Drug Dependence and Alcoholism (SSADDA),<sup>25, 26</sup> which yields DSM-IV and DSM-5 diagnoses of lifetime alcohol and drug dependence and other major psychiatric traits. Lifetime trauma exposure was defined as previous experience or witnessing of traumatic events including military combat; an assault, rape, or kidnapping; seeing someone seriously injured or killed; a flood, earthquake, large fire, or other disaster; an airplane crash or serious car accident; a shooting or bombing; or any situation where you feared there was a serious threat to your life or the life of another person. DSM-5 AUD criterion count was used as a dimensional measure based on the 11 DSM diagnostic criteria for AUD. Further details are reported in our previous study.<sup>15</sup> In the analysis, we included only subjects who reported having ever consumed alcohol (i.e., were alcohol exposed).

**Genetics**—The Illumina HumanOmni1-Quad v1.0 microarray was used to genotype 5,546 subjects (Yale-Penn1) at the Center for Inherited Disease Research or the Yale Center for Genome Analysis; and the Illumina HumanCoreExome array was used to genotype 2,538 additional subjects at the Gelernter Lab at Yale (Yale-Penn2). Genotyping, Imputation, ancestry assignment, and PC analysis are described in our previous articles.<sup>4, 21–24</sup>

### Data Analysis

A proxy-phenotype analysis was conducted considering the ASTARRS cohorts (ASTARRS1 and ASTARRS2) as the discovery sample (Stage-1) and the Yale-Penn cohorts (Yale-Penn1 and Yale-Penn2) as the replication sample (Stage-2). Proxy-phenotype analysis is a two-stage research strategy proposed by Rietveld and colleagues.<sup>15</sup> In the first stage, a proxy phenotype (i.e., alcohol misuse) was used to identify a relatively small set of SNPs considering a suggestive statistical significance threshold (i.e.,  $p < 5 \times 10^{-5}$ ). In the second

stage, this set of candidate SNPs was tested in an independent sample with respect to the phenotype of interest (i.e., AUD) at a significance threshold corrected for the number of proxy-associated SNPs. Consistent with the National Institute of Mental Health's Research Domain Criteria (RDoC) initiative,<sup>27–29</sup> we considered two-dimensional measures derived from symptom scales based on self-report information: an alcohol misuse score in the ASTARRS cohorts, and a DSM-5 AUD criterion count in the Yale-Penn cohorts. Dichotomous trauma exposure was considered as interactive factor, because it was available in both cohorts. A genome-wide gene-by-trauma interaction analysis was conducted in the ASTARRS cohorts (stage-1). Considering linkage disequilibrium (LD)-independent variants with  $p < 5 \times 10^{-5}$  in the ASTARRS analysis, a replication analysis was performed in the Yale-Penn cohorts (stage-2). Independent variants were defined as more than 500 kb distant and with  $r^2 < 0.2$ . SNPs that survived a Bonferroni correction that accounted for the number of independent loci were considered a replication. The statistical power of our GEWIS was calculated using QUANTO software (available at <http://biostats.usc.edu/Quanto.html>). In the ASTARRS sample, we have 92.7% statistical power to detect a moderate GxE effect ( $\beta_{G \times E} = 0.7$ ) at significance level  $p < 5 \times 10^{-5}$  for alleles with frequency 5%. In the Yale-Penn sample, we have 88% statistical power to detect a moderate GxE effect ( $\beta_{G \times E} = 0.7$ ) at significance level  $p < 5 \times 10^{-4}$  for alleles with frequency 5%.

Plink 1.9<sup>30</sup> was used to conduct the analysis in the ASTARRS cohort, and the interaction test was based on comparing the difference between regression coefficients in the trauma-exposed subjects vs. the trauma-unexposed subjects. Because the Yale-Penn cohorts include related individuals, we performed the interaction analysis using the R package GWAF<sup>31</sup> to fit a generalized estimating equations (GEE) model to adjust for correlations among related individuals. To verify that no important differences between the two methods were present, we tested the GWAF package and Plink 1.9 in a cohort of unrelated subjects and observed negligible differences due to number approximations. In both analyses, we included SNPs with minor allele frequency 5% and high imputation quality (Info 0.8). Before being entered into the analysis, alcohol misuse dimensional scales (i.e., alcohol misuse count and DSM-5 AUD criterion count) were adjusted for age, sex, and the top-10 ancestry PCs, and then normalized using appropriate Box-Cox power transformations. In every analysis, the samples were stratified by genotyping array and ancestry, and the results were combined by meta-analysis using the program METAL.<sup>32</sup> Functional annotation of the identified loci was conducted using HaploReg v4.1.<sup>33</sup>

Finally, we evaluated whether the number of loci that showed the same effect directions in the ASTARRS and the Yale-Penn cohorts was significantly different from chance, by conducting a permutation analysis; and then performed a gene ontology (GO)-enrichment analysis on the basis of the direction-replicated loci using DAVID 6.8 beta version (released May 2016; available at <https://david-d.ncicrf.gov/home.jsp>). Fisher's exact tests and Bonferroni multiple testing corrections were applied in the GO-enrichment analysis. We also used WebGestalt<sup>34</sup> (available at <http://www.webgestalt.org/>) to conduct an enrichment analysis for KEGG pathways using hypergeometric statistical method and Benjamini-Hochberg multiple test adjustment with significance level set at 0.05.

## Results

Table 1 reports the characteristics of the cohorts included in our GEWIS. Both ASTARRS and Yale-Penn samples include subjects genotyped with different platforms. Accordingly, we stratified the samples according to their genotyping platform and used a meta-analytic approach to integrate the results for Stage-1 (ASTARRS1 and ASTARRS2) and Stage-2 (Yale-Penn1 and Yale-Penn2) cohorts. Within each study population (ASTARRS and Yale-Penn), the cohorts have similar characteristics (age, sex, ancestry, and trauma exposure), except for sex between the ASTARRS cohorts (ASTARRS1 women = 12%; ASTARRS2 women = 22%). Considering the differences between the two study populations (ASTARRS vs. Yale-Penn), the ASTARRS cohorts are mainly constituted by young European-descent subjects, while the Yale-Penn cohorts mainly consist of older participants with both sexes and ancestries (European and African) almost equally represented. No Hispanic-American group is included in the Yale-Penn cohorts. Lifetime trauma exposure is slightly higher in the ASTARRS cohorts than that observed in the Yale-Penn cohorts (77% vs. 65%).

In the first stage, the genome-wide gene-by-trauma interaction analysis of alcohol misuse showed negligible inflation or deflation in the ancestry-stratified investigations of the ASTARRS cohorts (Supplemental Table 2). This confirmed that no systematic bias affected our GEWIS. GxE analysis can be biased by interactions among predictors and this can be hardly detected in a candidate gene study.<sup>35</sup> In a GEWIS a systematic bias in the GxE model would have caused inflation in the distribution of the test statistics.

There was no genome-wide significant result in the ancestry-specific or the trans-population meta-analyses. Considering  $p < 5 \times 10^{-5}$  as the significance threshold for follow-up in the second stage of the proxy-phenotype analysis, we identified 68, 49, and 45 independent loci in African-specific, European-specific, and trans-population meta-analyses, respectively (Supplemental Table 3). In African-American subjects, we identified an interaction of *PRKGI* rs1729578 with trauma exposure in the ASTARRS cohort and replicated its interaction with trauma exposure in the Yale-Penn cohort (discovery-replication meta-analysis:  $z=5.64$ ,  $p=1.69 \times 10^{-8}$ ; ASTARRS AA:  $z=4.46$ ,  $p=8.09 \times 10^{-6}$ ; Yale-Penn AA:  $z=3.62$ ,  $p=2.98 \times 10^{-4}$ ). Table 2 shows the details of the association of rs1729578 with alcohol misuse in trauma-exposed and unexposed subjects and its interaction with trauma exposure in relation to alcohol misuse. In the meta-analysis of the discovery (Stage-1) and replication (Stage-2) AA cohorts ( $N=6,744$ ), rs1729578 showed a genome-wide significant interaction with trauma-exposure in relation to alcohol misuse ( $z=5.64$ ,  $p=1.69 \times 10^{-8}$ ; Figure 1). Rs1729578\*C allele was positively associated with alcohol misuse in trauma-exposed subjects ( $z=4.27$ ,  $p=1.96 \times 10^{-5}$ ); and was negatively associated in trauma-unexposed subjects ( $z=-5.29$ ,  $p=1.21 \times 10^{-7}$ ). No significant replication was observed in the EA or trans-population meta-analysis. The Hispanic-specific meta-analysis could not be included in the second stage because no Hispanic group was available for replication in the Yale-Penn cohorts.

Rs1729578 is located in an intron of *PRKGI* (cGMP-dependent protein kinase 1) and in AAs it is in high LD ( $r^2 > 0.8$ ) with other two *PRKGI* intronic variants (rs1194520 and rs871995). According to the data provided by the Roadmap Epigenomics Consortium and

the ENCODE Project Consortium,<sup>36, 37</sup> rs1729578 is associated with a MIZF motif change; rs1194520 is located in enhancer histone marks (6 tissues), DNase sites (16 tissues), a CFOS protein bound site, and it is associated with 4 altered transcription factor (TF) motifs (Irf, SETDB1, STAT, YY1). Similarly, rs871995 is located in promoter histone marks (18 tissues), enhancer histone marks (8 tissues), DNase sites (1 tissue), and is associated with 3 TF altered motifs (Foxp1, Hoxa10, Irf). Details on chromatin-state annotation of the *PRKGI* variants are provided in Supplemental Table 4.

Finally, we verified whether the loci identified in Stage-1 had the same effect directions in the Stage-2 cohorts. In all of the analyses (African-specific, European-specific, and trans-population), we observed that the number of Stage1-identified loci with the same effect directions in the Stage-2 cohorts was higher than would be expected by chance (Figure 2): 42 loci in AAs ( $p_{\text{permutation}} = 0.022$ ); 33 loci in EAs ( $p_{\text{permutation}} = 0.004$ ); and 28 loci in the trans-population analysis ( $p_{\text{permutation}} = 0.041$ ). Considering these loci with the same direction in both cohorts (Supplemental Table 5), we observed significant enrichment for: GO:0016286~*small conductance calcium-activated potassium channel activity* in AAs; GO:0040011~*locomotion* in EAs; GO:0050890~*cognition* and GO:0042517~*positive regulation of tyrosine phosphorylation of Stat3 protein*. Details regarding the results of enrichment analysis are reported in Table 3. The analysis based on KEGG pathways identified several significant enrichments (Supplemental Table 6). Calcium signaling pathway (KEGG ID: 04020) was also confirmed by this analysis. Other relevant molecular pathways were: Cytokine-cytokine receptor interaction (KEGG ID: 04060); Long-term potentiation (KEGG ID: 04720); Insulin signaling pathway (KEGG ID: 04910).

## Discussion

To our knowledge, this study, which included 24,445 individuals, is the largest GEWIS in psychiatric genetics, and the first examining risk for alcohol misuse. Because the cohorts investigated were assessed using different instruments and criteria intended to measure similar domains, we applied the proxy-phenotype approach.<sup>16</sup>

In AAs, our proxy-phenotype analysis uncovered a variant, rs1729578, identified in the ASTARRS cohorts and replicated in the Yale-Penn samples. The rs1729578\*C allele showed an interaction with trauma exposure in relation to alcohol misuse symptoms. Although it has similar allele frequencies in European- and African-ancestry populations, no effect was observed in the European cohort, possibly due to a different haplotype structure in these groups.<sup>38</sup> In African populations, the variant is in high LD ( $r^2 > 0.8$ ) with other two variants, rs1194520 and rs871995. These SNPs are located in an intron of *PRKGI*, the gene encoding cGMP-dependent protein kinase 1. Functional annotation from the Roadmap Epigenomics Consortium and the ENCODE Project Consortium indicated that these variants are located in genomic regulatory regions.<sup>36, 37</sup> Specifically, they are located within multiple chromatin marks across different tissues. Previous studies have demonstrated that non-coding variants identified by GWAS are enriched for chromatin modifications.<sup>39</sup> The *PRKGI* protein product, cGMP-dependent protein kinase 1, corresponds to both the type I alpha and type I beta isoforms of cyclic guanosine monophosphate (cGMP)-dependent protein kinase, by alternative transcript splicing.<sup>40, 41</sup> The gene is most strongly expressed in



smooth muscle, platelets, cerebellar Purkinje cells, hippocampal neurons, and the lateral amygdala.<sup>40, 41</sup> cGMP plays an important role in learning and memory,<sup>42</sup> and a *PRKG1* knockout mouse model showed alterations in the cerebellar phospho-proteome that suggests impaired cerebellar long-term depression at Purkinje cell synapses.<sup>43</sup> *PRKG1* mutant mice also show differences from wild-type mice in circadian rhythms, sleep and distinct aspects of learning.<sup>44</sup> Mouse models also demonstrate that cGMP-dependent protein kinase 1 in the amygdala is critical for auditory-cued fear memory and long-term potentiation.<sup>45, 46</sup> *In vitro* studies also show that cGMP-dependent protein kinase 1 isoforms are involved in serotonin transporter regulation.<sup>47</sup> In *Drosophila melanogaster*, the homolog of the human *PRKG1*, the well-studied *foraging* (*for*) gene encodes a cGMP-dependent protein kinase (PKG). Two *for* variants have been observed in nature: rover allele (*for<sup>R</sup>*) with high PKG activity; and sitter allele (*for<sup>S</sup>*) with low PKG activity.<sup>48</sup> In *Drosophila*, PKG activity interacts with early life stress in determining adult exploratory and fitness traits.<sup>49, 50</sup> PKG activity seems to control synaptic transmission tolerance to acute stress at the *Drosophila* larval neuromuscular junction, where inhibition promotes functional protection, while activation increases susceptibility to neurotransmission breakdown.<sup>51</sup> A GWAS of post-traumatic stress disorder identified *PRKG1* as a risk locus in a military cohort independent from those investigated in the current studies,<sup>52</sup> also supporting the role of *PRKG1* in stress-response related traits in humans.

Finally, our proxy-phenotype analysis demonstrated that the enrichment for loci with same effect direction in stage-1 and stage-2 was unlikely to be due to chance. Investigating these loci, we observed indicative GO enrichments. The enrichment for GO:0016286~*small conductance calcium-activated potassium channel activity* confirmed previous findings of genome-wide genetic and epigenetic investigations on the role of potassium channels and calcium metabolism in substance use disorders.<sup>21, 22, 53</sup> Both GO:0040011~*locomotion* and GO:0050890~*cognition* appear consistent with the role of genes involved in mobility and cognitive functions highlighted by the *PRKG1* result. GO:0042517~*positive regulation of tyrosine phosphorylation of Stat3 protein* suggests that mechanisms related to *STAT3* regulation are involved in the trauma response. *STAT3* is involved in signal transduction and transcription activation of a wide range of genes in response to cell stimuli,<sup>54</sup> but no studies have yet investigated its involvement in the behavioral stress response. Pathway-based enrichment analysis confirmed the GO result related to the calcium metabolism (Calcium signaling pathway, KEGG ID: 04020). Besides this confirmatory result, the most significant pathway enrichment was observed for Cytokine-cytokine receptor interaction (KEGG ID: 04060). This is a relevant pathway related to immune functions such as inflammatory host defenses, cell growth, differentiation, cell death, angiogenesis, and development and repair processes aimed at the restoration of homeostasis. Our previous GWAS of PTSD showed a strong overlapping with autoimmune diseases<sup>19</sup> and our current finding confirm the role of immune functions in trauma response.

In conclusion, our study provides the first genome-wide evidence regarding a mechanism by which traumatic life events interact with human genetic variation in relation to alcohol misuse. However, although to our knowledge this is the largest GEWIS performed in psychiatric genetics, our current findings are still limited and further GEWIS with larger sample sizes together with translational studies will be needed to uncover more completely

how traumatic experience interacts with the genetic predisposition to modify risk for alcohol use disorders. Our main result is the identification of *PRKG1* rs1729578 as a risk locus interacting with trauma exposure in determining alcohol misuse. Our study design included different types of trauma within the “lifetime trauma exposure” category. Since certain traumatic experiences may be consequences of the alcohol abuse (e.g., individuals who abuse alcohol are more likely to have motor vehicle collisions than those who do not abuse alcohol), further studies are needed to dissect the complex interactive network of alcohol misuse, trauma experience, and genetics. Additional experiments are also necessary to understand the molecular mechanisms (e.g., epigenetic changes, transcriptional regulation) involved in this interactive process. However, the *PRKG1* locus encodes a prospective target for the development of pharmacological treatments for stress-response related traits. Indeed, an *in vivo* study has shown that the blockade of  $\alpha 1$ -adrenergic receptors mitigates stress-disturbed cGMP signaling.<sup>55</sup> Further studies should consider whether pharmacological treatments targeted at the cGMP signaling system are useful to reduce risk behaviors, such as alcohol misuse, in subjects exposed to traumatic events. Other potential targets for future translational investigations of trauma-related psychopathologies include potassium channels, calcium metabolism, and *STAT3* regulation system.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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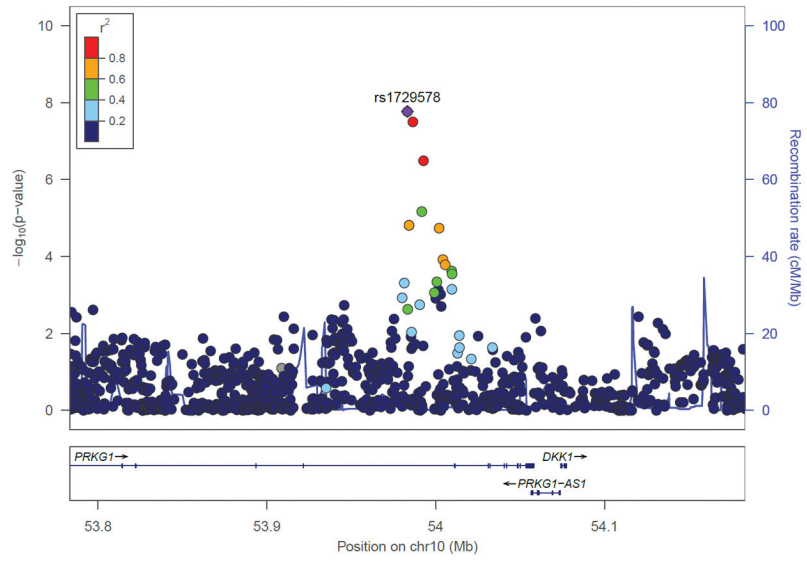
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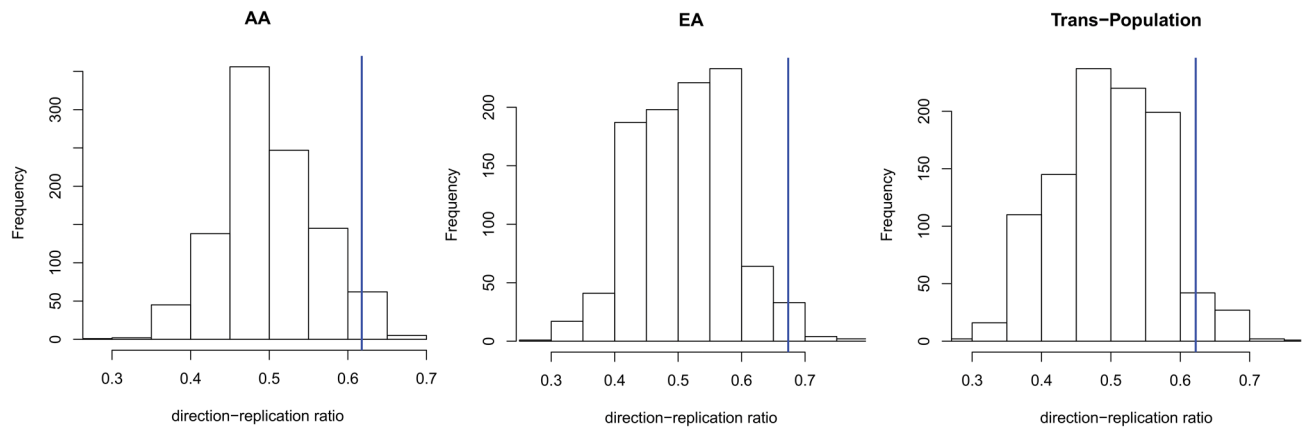
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**Figure 1.** Regional Manhattan Plots of *PRKG1* rs1729578 in African American overall meta-analysis (ASTARRS cohorts + Yale-Penn cohorts).



**Figure 2.** Null distribution of direction-replication ratios generated through 1,000 random permutations. Blue lines indicate observed values.

Characteristics of the ASTARRS and Yale-Penn cohorts investigated in the proxy-phenotype analysis.

**Table 1**

| Characteristics                  | ASTARRS1 (n = 14,000) | ASTARRS2 (n = 2,361) | Yale-Penn1 (n = 5,546) | Yale-Penn2 (n = 2,538) |
|----------------------------------|-----------------------|----------------------|------------------------|------------------------|
| Age, mean (SD)                   | 24 (5)                | 20 (3)               | 40 (10)                | 40 (12)                |
| Sex (women), n (%)               | 1,631 (12)            | 510 (22)             | 2,472 (45)             | 989 (39)               |
| African-Americans, n (%)         | 2,056 (15)            | 335 (14)             | 3,215 (58)             | 1,114 (44)             |
| European-Americans, n (%)        | 9,146 (65)            | 1,586 (67)           | 2,331 (42)             | 1,424 (56)             |
| Hispanic-Americans, n (%)        | 2,798 (20)            | 440 (19)             | -                      | -                      |
| Lifetime Trauma (exposed), n (%) | 10,643 (76)           | 1,874 (79)           | 3,552 (64)             | 1,694 (67)             |



Association of *PRKG1* rs1729578\*C allele with alcohol-related dimensional scales in subjects of African descent exposed and unexposed to lifetime trauma.

**Table 2**

| Cohort               | Allele Frequency | Lifetime Trauma |        |                       |                       | No Lifetime Trauma |                       |         |                       | Interaction |      |                       |  |
|----------------------|------------------|-----------------|--------|-----------------------|-----------------------|--------------------|-----------------------|---------|-----------------------|-------------|------|-----------------------|--|
|                      |                  | n               | Beta   | SE                    | P value               | n                  | Beta                  | SE      | P value               | Beta        | SE   | P value               |  |
| <i>ASTARRS1</i>      | 0.25             | 1,483           | 0.05   | 0.03                  | 0.081                 | 573                | -0.19                 | 0.05    | 1.05*10 <sup>-4</sup> | 0.25        | 0.06 | 1.15*10 <sup>-5</sup> |  |
| <i>ASTARRS2</i>      | 0.21             | 265             | -0.02  | 0.09                  | 0.775                 | 70                 | -0.25                 | 0.13    | 0.068                 | 0.22        | 0.16 | 0.161                 |  |
| <i>Yale-Penn1</i>    | 0.23             | 1,986           | 0.20   | 0.07                  | 3.49*10 <sup>-3</sup> | 1,229              | -0.21                 | 0.08    | 7.79*10 <sup>-3</sup> | 0.10        | 0.03 | 8.92*10 <sup>-5</sup> |  |
| <i>Yale-Penn2</i>    | 0.22             | 752             | -0.02  | 0.13                  | 0.846                 | 362                | -0.12                 | 0.16    | 0.436                 | 0.02        | 0.05 | 0.625                 |  |
| <i>Meta-analysis</i> | 0.23             | 4,486           | Z=4.27 | 1.96*10 <sup>-5</sup> | 2,234                 | Z=-5.29            | 1.21*10 <sup>-7</sup> | Z=-5.64 | 1.69*10 <sup>-8</sup> |             |      |                       |  |

Results from the enrichment analysis conducted considering the loci with same direction in discovery and replication cohorts in trans-population, AA, and EA analyses.

**Table 3**

| Analysis         | GO term   | Genes                           | Fisher's exact P value | Bonferroni-corrected P value |
|------------------|---|---------------------------------|------------------------|------------------------------|
| Trans-population | GO:0050890~cognition  | <i>CBR3, CTNS, DOPEY2, GRM5</i> | $1.9 \times 10^{-6}$   | $7.1 \times 10^{-4}$         |
|                  | GO:0042517~positive regulation of tyrosine phosphorylation of Stat3 protein | <i>CLCF1, IL23R, VEGFA</i>      | $6.4 \times 10^{-5}$   | $1.6 \times 10^{-2}$         |
| AA               | GO:0016286~small conductance calcium-activated potassium channel activity   | <i>KCNN2, KCNN3</i>             | $2.3 \times 10^{-5}$   | $5.4 \times 10^{-3}$         |
| EA               | GO:0040011~locomotion   | <i>ATP2B2, JPH3</i>             | $6.7 \times 10^{-5}$   | $5.4 \times 10^{-3}$         |