



Epidemiology Faculty Publications

Epidemiology

11-15-2016

A DNA Methylation Biomarker of Alcohol Consumption

C. Liu The Framingham Heart Study

R. E. Marioni *University of Edinburgh, UK*

Å. K. Hedman Uppsala University, Sweden

L. Pfeiffer German Research Center for Environmental Health, Germany

P. -C. Tsai King's College London, UK

See next page for additional authors

Right click to open a feedback form in a new tab to let us know how this document benefits you.

Follow this and additional works at: https://uknowledge.uky.edu/epidemiology_facpub Part of the <u>Epidemiology Commons</u>, <u>Genetics and Genomics Commons</u>, and the <u>Psychiatry</u> Commons

Repository Citation

Liu, C.; Marioni, R. E.; Hedman, Å. K.; Pfeiffer, L.; Tsai, P. -C.; Reynolds, L. M.; Just, A. C.; Duan, Q.; Boer, C. G.; Tanaka, T.; Elks, C. E.; Aslibekyan, S.; Brody, J. A.; Kühnel, B.; Herder, C.; Almli, L. M.; Zhi, D.; Wang, Y.; Huan, T.; Yao, C.; Mendelson, M. M.; Joehanes, R.; Liang, L.; Love, S. -A.; Guan, W.; Shah, S.; McRae, A. F.; Kretschmer, A.; Prokisch, H.; Strauch, K.; and Arnett, Donna K., "A DNA Methylation Biomarker of Alcohol Consumption" (2016). *Epidemiology Faculty Publications*. 35. https://uknowledge.uky.edu/epidemiology_facub/35

This Article is brought to you for free and open access by the Epidemiology at UKnowledge. It has been accepted for inclusion in Epidemiology Faculty Publications by an authorized administrator of UKnowledge. For more information, please contact UKnowledge@lsv.uky.edu.

Authors

C. Liu, R. E. Marioni, Å. K. Hedman, L. Pfeiffer, P. -C. Tsai, L. M. Reynolds, A. C. Just, Q. Duan, C. G. Boer, T. Tanaka, C. E. Elks, S. Aslibekyan, J. A. Brody, B. Kühnel, C. Herder, L. M. Almli, D. Zhi, Y. Wang, T. Huan, C. Yao, M. M. Mendelson, R. Joehanes, L. Liang, S. -A. Love, W. Guan, S. Shah, A. F. McRae, A. Kretschmer, H. Prokisch, K. Strauch, and Donna K. Arnett

A DNA Methylation Biomarker of Alcohol Consumption

Notes/Citation Information

Published in Molecular Psychiatry, v. 23, issue 2, p. 422-433.

© The Author(s) 2018

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/4.0/.

Due to the large number of authors, only the first 30 and the authors affiliated with the University of Kentucky are listed in the author section above. For the complete list of authors, please download this article or visit: https://doi.org/10.1038/mp.2016.192

Digital Object Identifier (DOI) https://doi.org/10.1038/mp.2016.192

ORIGINAL ARTICLE A DNA methylation biomarker of alcohol consumption

C Liu^{1,2,3,55}, RE Marioni^{4,5,6,55}, ÅK Hedman^{7,55}, L Pfeiffer^{8,9,55}, P-C Tsai^{10,55}, LM Reynolds^{11,55}, AC Just^{12,55}, Q Duan^{13,55}, CG Boer^{14,55}, T Tanaka^{15,55}, CE Elks¹⁶, S Aslibekyan¹⁷, JA Brody¹⁸, B Kühnel^{8,9}, C Herder^{19,20}, LM Almli²¹, D Zhi²², Y Wang²³, T Huan^{1,2}, C Yao^{1,2}, MM Mendelson^{1,2}, R Joehanes^{1,2,24}, L Liang²⁵, S-A Love²³, W Guan²⁶, S Shah^{6,27}, AF McRae^{6,27}, A Kretschmer^{8,9}, H Prokisch^{28,29}, K Strauch^{30,31}, A Peters^{8,9,32}, PM Visscher^{4,6,27}, NR Wray^{6,27}, X Guo³³, KL Wiggins¹⁸, AK Smith²¹, EB Binder³⁴, KJ Ressler³⁵, MR Irvin¹⁷, DM Absher³⁶, D Hernandez³⁷, L Ferrucci¹⁵, S Bandinelli³⁸, K Lohman¹¹, J Ding³⁹, L Trevisi⁴⁰, S Gustafsson⁷, JH Sandling^{41,42}, L Stolk¹⁴, AG Uitterlinden^{14,43}, I Yet¹⁰, JE Castillo-Fernandez¹⁰, TD Spector¹⁰, JD Schwartz⁴⁴, P Vokonas⁴⁵, L Lind⁴⁶, Y Li⁴⁷, M Fornage⁴⁸, DK Arnett⁴⁹, NJ Wareham¹⁶, N Sotoodehnia¹⁸, KK Ong¹⁶, JBJ van Meurs¹⁴, KN Conneely⁵⁰, AA Baccarelli⁵¹, IJ Deary^{4,52}, JT Bell¹⁰, KE North^{23,56}, Y Liu^{11,56}, M Waldenberger^{8,9,56}, SJ London^{53,56}, E Ingelsson^{7,54,56} and D Levy^{1,2,56}

The lack of reliable measures of alcohol intake is a major obstacle to the diagnosis and treatment of alcohol-related diseases. Epigenetic modifications such as DNA methylation may provide novel biomarkers of alcohol use. To examine this possibility, we performed an epigenome-wide association study of methylation of cytosine-phosphate-guanine dinucleotide (CpG) sites in relation to alcohol intake in 13 population-based cohorts ($n_{total} = 13 317$; 54% women; mean age across cohorts 42–76 years) using whole blood (9643 European and 2423 African ancestries) or monocyte-derived DNA (588 European, 263 African and 400 Hispanic ancestry) samples. We performed meta-analysis and variable selection in whole-blood samples of people of European ancestry (n = 6926) and identified 144 CpGs that provided substantial discrimination (area under the curve = 0.90–0.99) for current heavy alcohol intake (≥ 42 g per day in men and ≥ 28 g per day in women) in four replication cohorts. The ancestry-stratified meta-analysis in whole blood identified 328 (9643 European ancestry samples) and 165 (2423 African ancestry samples) alcohol-related CpGs at Bonferroni-adjusted $P < 1 \times 10^{-7}$. Analysis of the monocyte-derived DNA (n = 1251) identified 62 alcohol-related CpGs at $P < 1 \times 10^{-7}$. In whole-blood samples of people of European ancestry, we detected differential methylation in two neurotransmitter receptor genes, the γ -Aminobutyric acid-A receptor delta and γ -aminobutyric acid B receptor subunit 1; their

¹The Framingham Heart Study, Framingham, MA, USA; ²The Population Sciences Branch, Division of Intramural Research, National Heart, Lung and Blood Institute, Bethesda, MD, USA; ³Department of Biostatistics, Boston University School of Public Health, Boston, MA, USA; ⁴Centre for Cognitive Ageing and Cognitive Epidemiology, University of Edinburgh, Edinburgh, UK; ⁵Medical Genetics Section, Centre for Genomic and Experimental Medicine, Institute of Genetics and Molecular Medicine, University of Edinburgh, Edinburgh, UK; ⁶Queensland Brain Institute, The University of Queensland, Brisbane, QLD, Australia; ⁷Department of Medical Sciences, Molecular Epidemiology and Science for Life Laboratory, Uppsala University, Uppsala, Sweden; ⁸Research Unit of Molecular Epidemiology, Helmholtz Zentrum München, German Research Center for Environmental Health, Neuherberg, Germany; ⁹Institute of Epidemiology II, Helmholtz Zentrum München, German Research Center for Environmental Health, Neuherberg, Germany; ¹⁰Department of Twin Research and Genetic Epidemiology, King's College London, London, UK; ¹¹Division of Public Health Sciences, Wake Forest School of Medicine, Winston-Salem, NC, USA; ¹²Department of Preventive Medicine, Icahn School of Medicine at Mount Sinai, New York, NY, USA; ¹³Department of Genetics, University of North Carolina, Chapel Hill, NC, USA; ¹⁴Department of Internal Medicine, Erasmus MC, Rotterdam, The Netherlands; ¹⁵Translational Gerontology Branch, National Institute on Aging, Baltimore, MD, USA; ¹⁶MRC Epidemiology Unit, Institute of Metabolic Science, University of Cambridge, Cambridge, UK; ¹⁷Department of Epidemiology, University of Alabama at Birmingham, Birmingham, AL, USA; ¹⁸Cardiovascular Health Research Unit, Department of Medicine, University of Washington, Seattle, WA, USA; ¹⁹German Center for Diabetes Research (DZD), München-Neuherberg, Germany; ²⁰Institute for Clinical Diabetology, German Diabetes Center, Leibniz Center for Diabetes Research at Heinrich Heine University, Düsseldorf, Germany; ²¹Department of Psychiatry and Behavioral Sciences, Emory University School of Medicine, Atlanta, GA, USA; ²²School of Biomedical Informatics and School of Public Health, The University of Texas Health Science Center at Houston, Houston, TX, USA; ²³Department of Epidemiology, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA; ²⁴Hebrew SeniorLife, Harvard Medical School, Boston, MA, USA; ²⁵Harvard School of Public Health, Harvard University, Boston, MA, USA; ²⁶Division of Biostatistics, School of Public Health, University of Minnesota, Minneapolis, MN, USA; 27 Institute for Molecular Bioscience, The University of Queensland, Brisbane, QLD, Australia; 28 Institute of Human Genetics, Helmholtz Zentrum München, German Research Center for Environmental Health, Neuherberg, Germany; ²⁹Institute of Human Genetics, Technische Universität München, München, Germany; ³⁰Institute of Medical Informatics, Biometry and Epidemiology, Ludwig-Maximilians-Universität, Munich, Germany; ³¹Institute of Genetic Epidemiology, Helmholtz Zentrum München - German Research Center for Environmental Health, Neuherberg, Germany; ³²DZHK (German Centre for Cardiovascular Research), partner site Munich Heart Alliance, Munich, Germany; ³³Institute for Translational Genomics and Population Sciences, Department of Pediatrics, LABioMed at Harbor-UCLA Medical Center, Torrance, CA, USA; ³⁴Max-Planck Institute of Psychiatry, Munich, Germany; ³⁵Division of Depression and Anxiety Disorders, McLean Hospital, Belmont, MA, USA; ³⁶HudsonAlpha Institute for Biotechnology, Huntsville, AL, USA; ³⁷Laboratory of Neurogenetics, National Institute on Aging, National Institutes of Health, Bethesda, MD, USA; ³⁸Geriatric Unit, Azienda Sanitaria Firenze (ASF), Florence, Italy; ³⁹Department of Internal Medicine, Wake Forest School of Medicine, Winston-Salem, NC, USA; ⁴⁰Department of Environmental Health, Harvard T.H. Chan School of Public Health, Boston, MA, USA; ⁴¹Department of Medical Sciences, Rheumatology and Science for Life Laboratory, Uppsala University, Uppsala, Sweden; ⁴²Department of Medical Sciences, Molecular Medicine and Science for Life Laboratory, Uppsala University, Uppsala, Sweden; ⁴³Department of Epidemiology, Erasmus MC, Rotterdam, The Netherlands; ⁴⁴Departments of Environmental Health and Epidemiology, Harvard T.H. Chan School of Public Health, Boston, MA, USA; ⁴⁵VA Boston Healthcare System and Boston University Schools of Public Health and Medicine, Jamaica Plain, MA, USA; ⁴⁶Department of Medical Sciences, Cardiovascular Epidemiology, Uppsala University, Uppsala, Sweden; 47 Department of Genetics, Department of Biostatistics and Department of Computer Science, University of North Carolina, Chapel Hill, NC, USA; 48 Institute of Molecular Medicine and Human Genetics Center, University of Texas Health Science Center at Houston, Houston TX, USA; 49 College of Public Health, University of Kentucky, Lexington, KY, USA; ⁵⁰Department of Human Genetics, Emory University School of Medicine, Atlanta, GA, USA; ⁵¹Department of Environmental Health Sciences, Mailman School of Public Health, Columbia University, New York, NY, USA; 52Department of Psychology, University of Edinburgh, Edinburgh, UK; 53Department of Health and Human Services, Division of Intramural Research, National Institute of Environmental Health Sciences, National Institutes of Health, Research Triangle Park, NC, USA and ⁵⁴Department of Medicine, Division of Cardiovascular Medicine, Stanford University School of Medicine, Stanford, CA, USA. Correspondence: Dr C Liu or D Levy, The Population Sciences Branch, Division of Intramural Research, The Framingham Heart Study, National Heart, Lung and Blood Institute, Perini Building, Framingham, MA 01701, USA. E-mail: chunyu.liu@nih.gov or levyd@nhlbi.nih.gov

⁵⁵These authors contributed equally to this work.

⁵⁶These authors jointly supervised the work.

Received 11 February 2016; revised 5 September 2016; accepted 14 September 2016; published online 15 November 2016

differential methylation was associated with expression levels of a number of genes involved in immune function. In conclusion, we have identified a robust alcohol-related DNA methylation signature and shown the potential utility of DNA methylation as a clinically useful diagnostic test to detect current heavy alcohol consumption.

Molecular Psychiatry (2018) 23, 422-433; doi:10.1038/mp.2016.192; published online 15 November 2016

INTRODUCTION

Each year, nearly 2.5 million deaths worldwide are attributable to alcohol use.¹ Most alcohol-attributable diseases and injuries occur in people without a diagnosed alcohol use disorder.²⁻⁵ Researchers have attempted to develop laboratory tests to detect heavy drinkers who are more reliable than self-reported alcohol intake (e.g. alcohol screening guestionnaires). In addition, a biomarker would be useful in epidemiologic studies of health effects of alcohol as an objective measure to supplement and validate selfreported data. It could also prove useful in studies of other exposures where careful adjustment for alcohol intake is needed.⁶ Several biochemical measurements, such as serum alanine transaminase (ALT) and aspartate transaminase (AST) levels have been used to assess heavy alcohol use. However, the discriminatory ability of these biomarkers is far from ideal, with the area under the curve (AUC) to predict heavy alcohol consumption ranging from 0.21 to 0.67.7 The addition of four protein markers, AT-rich interactive domain-containing protein 4B (ARID4B), phosphatidylcholine-sterol acyltransferase (LCAT), hepatocyte growth factor-like protein (MST1) and ADP-ribosylation factor 6 (ARL6), improved AUC values for the detection of heavy drinkers to 0.73–0.86, leaving room for further improvement.⁷

Emerging evidence suggests that alcohol consumption influences epigenetic modifications,^{8–10} which in turn can affect gene expression levels.^{8,11,12} Methylation of the cytosine position in CpGs is among the best-characterized epigenetic modifications.¹³ To date, more than 20 studies have been conducted to identify alcohol-related DNA methylation signatures. Most of these studies, however, have focused on alcohol dependence in relation to 'global' methylation levels or preselected candidate genes¹⁴ and only a few studies have used epigenome-wide approaches.15-18 The largest genome-wide study so far included about 700 individuals.¹⁶ To date, limited sample sizes have hindered the search for a robust alcohol-related DNA methylation signature. Hence, there is a need for a large-scale collaborative effort to determine the association of alcohol consumption with DNA methylation across the genome. Here we demonstrate that DNA methylation can be used as a highly predictive blood biomarker to detect heavy alcohol drinking. We also report meta-analysis results from epigenome-wide association studies (EWAS) in up to 13 317 individuals from 13 cohorts in which DNA methylation was measured in blood samples using the Infinium HumanMethylation450 BeadChip (Illumina, San Diego, CA, USA). Third, we determined genetic contributions to alcohol-related methylation the differences. Finally, we explored the functional implications of alcohol-related differential methylation by testing its association with gene expression in blood.

MATERIALS AND METHODS

Study population

This analysis included 13 317 participants from 13 population-based prospective cohorts of the Cohorts for Heart and Aging Research in Genomic Epidemiology Consortium plus (CHARGE+) Consortium. These cohorts were sampled from free-living members of the community, but they were all not required to be healthy nor were they selected based on disease. During follow-up, some participants developed health conditions such as cardiovascular diseases (CVDs) and cancer. About 54% of participants were women and the average age was from 42 to 76 years

old across the cohorts (Table 1). The patterns of alcohol consumption varied widely across the cohorts. For nine cohorts, fewer than one-third of participants reported no current alcohol intake and for four cohorts more than 50% of participants reported no current alcohol intake. The high proportion of non-drinkers in these four cohorts is in line with other studies of people of comparable age, birth cohort and gender mix. Heavy drinkers, defined below, represented 2–17% of participants across studies (Table 1). Informed consent for genetic studies was obtained from all subjects. The protocol for each study was approved by the institutional review board of each cohort.

Alcohol traits

Alcohol consumption was measured by self-administered questionnaires or structured interview with a trained psychologist at the same period when blood samples were obtained for DNA methylation quantification. Alcohol consumption measured the total consumption of beer, wine and spirits. For American cohorts, a drink was defined as 12 ounces of beer, 4-5 ounces of wine or 1.5 ounces of liquor, where one drink is equivalent to ~14 g of ethanol. For European cohorts, a slightly different definition of 'a drink' and its conversion to grams of ethanol was used (Supplementary Information: pp 14-16, 19-22 in Description of study samples). The continuous exposure variable was defined as the average grams of ethanol consumed per day (g per day) over the course of a year during the period when the blood sample was collected for DNA methylation quantification. The continuous variable was further categorized into four drinking categories. 'Non-drinkers' were subjects with no alcohol consumption (i.e., q per day = 0); 'light drinkers' were subjects who consumed 0 < qper day ≤ 28 in men and 0 < q per day ≤ 14 in women; 'at risk-drinkers' were subjects who consumed 28 < g per day < 42 in men and 14 < gper day < 28 in women; 'heavy drinkers' were subjects who consumed \geq 42 g per day in men and \geq 28 g per day in women.

To explore the effects of 'former' alcohol drinking on DNA methylation, we examined alcohol consumption at prior examinations for all current non-drinkers (n = 693, see Table 1) in the Framingham Heart Study (FHS) because information on prior drinking was not available in the majority of other cohorts. We classified non-drinkers in FHS into 'never' drinkers and 'former' drinkers. 'Never' drinkers were individuals who self-reported no alcohol consumption at any prior examination; 'former' drinkers were individuals who reported alcohol consumption at any prior examination. For 'former' drinkers, we calculated their alcohol consumption ('g per day') at each prior examination.

DNA methylation quantification and quality control

DNA was extracted from whole-blood (n = 9643 European (EA) and 2423 African ancestry (AA)) and CD14+ monocyte (n = 1251 of mixed EA (n = 588), AA (n = 263) and Hispanic ancestry (n = 400) samples (Table 1 and Supplementary Information). Detailed information about DNA extraction, bisulfite conversion, methylation profiling, normalization and quality control (QC) procedures can be found in Supplementary Information. Study samples were excluded from analysis if they had a missing rate of >1–5% across methylation probles; poor single nucleotide polymorphism (SNP) matching compared with previous genotyping of the 65 SNPs included on the methylation array; or sample outliers identified by multidimensional scaling techniques. The methylation probes were excluded if they were the 65 SNP probes, or probes that were previously identified to map to multiple locations (n = 29233);¹⁹ had average detection P > 0.01 (the detection *P*-value indicates the probe performance); had an underlying SNP within 10 bp of that probe or if the minor allele frequency (MAF) of the underlying SNP was >5% in the 1000 Genomes Project data ($n = 15 \ 178$).¹⁹ After these filtering procedures, ~ 440 000 DNA methylation probes remained for subsequent analyses.

Table 1. Characteristics of the study participants

Study	Ν	Men (%)	Age (years), mean (s.d.)	BMI mean (s.d.)	Current smoking (%)	g per day, Median (min, max)	Non-drinkers (%)	Light drinkers (%)	At-risk drinkers (%)	Heavy drinkers (%)
European ancest	ry whol	e blood (n=	9643)							
CHS	185	84 (45)	76 (5)	27 (5)	16 (9)	0 (0, 99)	104 (56)	59 (32)	7 (4)	15 (8)
EPIC-Norfolk	1275	650 (51)	60 (9)	27 (4)	191 (15)	3 (0, 98)	271 (21)	865 (68)	79 (6)	60 (5)
FHS	2427	1095 (45)	66 (9)	28 (5)	304 (13)	4 (0, 181)	693 (29)	1260 (52)	280 (11)	194 (8)
InCHIANTI	499	225 (45)	63 (16)	27 (4)	94 (19)	8 (0, 161)	106 (21)	265 (53)	70 (14)	58 (12)
KORA F4	1797	877 (49)	60 (9)	28 (5)	262 (15)	7 (0, 150)	534 (29)	751 (42)	282 (16)	230 (13)
LBC1936	920	465 (51)	70 (1)	28 (4)	103 (11)	7 (0,158)	181 (20)	574 (62)	104 (11)	61 (7)
NAS	623	623 (100)	72 (7)	28 (4)	27 (4)	6 (0, 93)	148 (24)	385 (62)	52 (8)	38 (6)
PIVUS	818	412 (50)	70 (0.2)	27 (4)	75 (9.2)	6.7 (0, 61)	142 (17)	639 (78)	32 (4)	5 (1)
RS	502	241 (48)	58 (7)	27 (5)	137 (27)	14 (0, 88)	10 (2)	366 (73)	84 (17)	42 (8)
TwinsUK	597	0 (0)	56 (9)	27 (5)	57 (10)	2 (0,59)	189 (31)	375 (63)	22 (4)	11 (2)
African ancestry	whole b	olood (n=24	23)							
ARIC	2003	721 (36)	56 (6)	30 (6)	490 (24)	0 (0, 301)	1519 (76)	67 (3)	69 (3)	348 (17)
CHS	190	66 (35)	73 (5)	29 (5)	29 (15.3)	0 (0, 74)	123 (65)	61(32)	2 (1)	4 (2)
GTP	230	76 (33)	42 (12)	32 (8)	74 (39)	14 (0,143)	45 (20)	113 (49)	NA	72 (31)
CD14+ monocyt	es (n = 1	251)								
MESA	1251	606 (48)	60 (9)	30 (6)	114 (9%)	8 (0, 191)	691 (55)	444 (36)	65 (5)	51 (4)

Abbreviations: ARIC, The Atherosclerosis Risk in Communities study; BMI, body mass index; CHS, The Cardiovascular Health Study; EPIC-Norfolk, The European Prospective Investigation into Cancer-Norfolk study; FHS, The Framingham Heart Study; GTP, The Grady Trauma Project; KORA F4, The Cooperative Health Research in the Region of Augsburg study; InCHIANTI, Invecchiare in Chianti; LBC1936, The Lothian Birth Cohort 1936; MESA, The Multi-Ethnic Study of Atherosclerosis; NAS, The Normative Aging Study; PIVUS, The Prospective Investigation of the Vasculature in Uppsala Seniors Study; RS, The Rotterdam Study; TwinsUK, The TwinsUK Study. The drinking categories were defined based on grams of alcohol consumed per day (g per day): non-drinkers, g per day = 0; light drinkers, $0 < - \leq 28$ g per day in men and $14 < - \leq 28$ g per day in women; and heavy drinkers, g per day ≥ 42 in men and ≥ 28 in women. The Monocyte samples included mixed samples of European (47%), African (21%) and Hispanic (32%) ancestries.

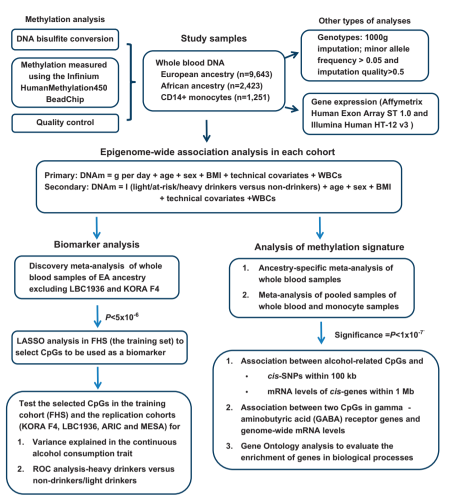


Figure 1. Overview of the study design. ARIC, The Atherosclerosis Risk in Communities study; BMI, body mass index; DNAm, DNA methylation value; FHS, the Framingham Heart Study; I (light/at-risk/heavy drinkers versus non-drinkers), the indicator variable for light drinkers versus non-drinkers, at-risk drinkers versus non-drinkers and heavy drinkers versus non-drinkers; KORA F4, The Cooperative Health Research in the Region of Augsburg study; LASSO, least absolute shrinkage and selection operator; LBC, The Lothian Birth Cohort; MESA, The Multi-Ethnic Study of Atherosclerosis; WBCs, white blood cell counts.

Clinical and laboratory phenotypes

Age, height and weight were measured using standard protocols implemented at the time DNA samples were collected. Body mass index (BMI) was calculated as weight (kg) per height (m) squared. Smoking was determined by self-report. Current smokers were defined as smokers of at least 1 cigarette per day over the course of a year at the time of blood sample collection for methylation quantification.

In the FHS cohort, the serum AST and ALT levels were measured on fasting morning samples using the kinetic method²⁰ at the same examination cycle when whole blood was obtained for DNA methylation measurement. ALT and AST were set to 5 if their measured levels were $< 5 \text{ U I}^{-1}$. An individual was defined as having CVD if he or she had coronary heart disease, myocardial infarction, atherothrombotic infarction of brain or congestive heart failure. A cancer phenotype was defined if the person had any type of cancer. Both CVD and cancer phenotypes were defined at the time of blood collection for measuring DNA methylation.

Statistical analysis

Epigenome-wide association studies. In each cohort, the primary EWAS model used a DNA methylation β -value (the ratio of methylated probe intensity divided by the sum of the methylation and unmethylated probe intensity) as the outcome variable and the continuous alcohol trait (g per day) as the predictor variable of interest. Because it has been shown that many CpG sites are significantly associated with age,²¹ sex²² and

BMI,²³ these three variables were adjusted for in EWAS analysis. Furthermore, it has been known that DNA methylation profiling is susceptible to batch effects²⁴ and by white cell composition in whole blood.²⁵ Thus, 'batch effects' and 'white cell blood counts' were adjusted for to minimize confounding effects that may result in spurious association. To account for batch effects, 'lab', 'experiment date', 'plate', 'row' and 'column' were adjusted for in the analyses. The measured or imputed²⁵ white blood cell counts for CD4 cells, CD8 cells, natural killer cells, B cells and monocytes were adjusted for in the analyses. Surrogate variables (to account for unknown confounders)²⁶ or principal components (estimated from genotypes to account for population stratification)²⁷were included in the EWAS model when applicable (pp 10, 11 and 15 of Description of study samples in Supplementary Information). In the secondary analysis, we used the categorical alcohol intake as the predictor adjusting for the same covariates described above. The non-drinker category was used as the reference group. We used a linear model in unrelated individuals or a linear mixed model in family samples to account for familial correlation in the association of DNA methylation and alcohol consumption.

It is unclear if smoking confounds the relationship between alcohol intake and DNA methylation or if smoking and alcohol intake are associated with common CpGs. Therefore, we performed an additional sensitivity analysis with and without current smoking status as a covariate in drinker-only samples. We compared the change in the regression coefficient for the continuous alcohol intake trait when including current 426

smoking in the model compared with the regression coefficient when smoking was not included in the model using the following equation:

$$\Delta \beta_{\text{alcohol}-\text{smk}} = 100\% * (\beta_{\text{alcohol}-\text{smk}} - \beta_{\text{alcohol}-\text{no}-\text{smk}})$$

We performed sensitivity tests in the FHS data to investigate if CVD and cancer confounded the relationship between DNA methylation and alcohol consumption. The sensitivity test compared the regression coefficients and *P*-values between methylation and alcohol intake in a EWAS model that did not adjust for CVD or cancer status to a second model that adjusted for CVD or cancer status. Other covariates included age, sex, BMI, batch effects and white blood cell counts.

DNA methylation as a biomarker in predicting alcohol consumption

We performed the following four-step analyses to investigate if DNA methylation can be used as a biomarker in discriminating alcohol consumption categories (Figure 1).

Step 1: To establish independent replication cohorts, we split the wholeblood DNA samples from 10 cohorts of (EA (n = 9643; Table 1) into separate discovery and replication sets. The discovery set consisted of eight EA cohorts (n = 6926), excluding the Lothian Birth Cohort 1936 (LBC1936) and Cooperative Health Research in the Region of Augsburg (KORA F4) study (Table 1). We performed a meta-analysis in the eight EA cohorts using an inverse-variance weighted random-effects model and selected CpGs at a relaxed threshold $P < 5 \times 10^{-6}$.

Step 2: To minimize overfitting and to explore which CpGs are more important for including in a biomarker of alcohol consumption, we performed least absolute shrinkage and selection operator (LASSO) regression in the FHS cohort as a training set:

$$\log\left(\text{g per day}+1\right) = \sum_{n=1} \text{ResidCpG} + \text{age} + \text{sex} + \text{BMI}$$

In the above formula, all CpGs at $P < 5 \times 10^{-6}$ were included simultaneously in the LASSO analysis. Because alcohol consumption was right skewed and contained non-drinkers, the log-transformed alcohol consumption (log (g per day+1)) was used as the outcome. To minimize potential confounding effects in selecting a set of CpGs as a biomarker, we obtained the residuals for each CpG in a linear regression model (CpG=age+sex+BMI+batch effects+white blood cell counts). Here variables for 'batch effects' and 'white blood cell counts' were the same as the variables used in EWAS analysis. In the LASSO analysis, we selected four sets of CpGs using *s*='lambda.min', 'lambda.1se', 0.08 and 0.12. The criterion *s*='lambda.min' selected the largest number of CpGs if they are not on the Infinium MethylationEPIC BeadChip (Illumina), which will replace the Illumina Infinium HumanMethylation450 BeadChip.

Step 3: In the FHS training set, we first estimated the proportion of variance in continuous alcohol consumption explained by the selected CpGs. The adjusted R^2 was estimated for the 'Null' model log (g per day + 1) = age + sex + BMI and the 'Full' model log (g per day + 1) = $\sum_{n=1}^{\infty} \text{ResidCpG} + \text{age} + \text{sex} + \text{BMI}$. The proportion of variance explained by a

set of CpGs was the difference of adjusted R^2 between the 'Full' and 'Null' models: $adjR_{CpGs}^2 = adjR_{age+sex+BMI+CpGs}^2 - adjR_{age+sex+BMI}^2$ Discrimination of heavy alcohol consumption from non-drinkers or light drinkers was our main focus. Therefore, we generated receiver-operating characteristic curves (ROC) in the FHS training cohort to evaluate the performance of these four sets of CpGs in classifying current heavy drinkers versus (1) non-drinkers, (2) light drinkers and (3) pooled individuals of light or non-drinkers. In addition, we evaluated if these CpGs can be used in classifying individuals in the following comparisons: (4) heavy drinkers versus at-risk-drinkers; (5) at-risk drinkers versus nondrinkers; (6) at-risk drinkers versus light drinkers; and (7) light drinkers versus non-drinkers. In all comparison pairs, the former category was the 'disease' group and the latter was the 'control' group. In ROC analysis, the expected probability of being 'diseased' was calculated using logistic regression in which the 'disease' (1/0) was used as the outcome variable, and age, sex and BMI without or with a set of CpGs (residuals) as independent variables. Sensitivity, specificity and the AUC for classifying 'diseased' individuals versus 'controls' were calculated. We also performed sensitivity tests to investigate the prediction performance from current smoking, ALT and AST.

Step 4: We repeated the Step 3 analyses in two independent cohorts of whole-blood-derived DNA samples in people of EA (LBC1936 and KORA F4)

for replication purposes. We also repeated the Step 3 analyses in wholeblood-derived DNA samples of people of AA (the Atherosclerosis Risk in Communities Study or ARIC) and in the monocyte-derived DNA samples (the Multi-Ethnic Study of Atherosclerosis or MESA) for both replication and generalization. The MESA samples included individuals of EA (n = 588), AA (n = 263) and Hispanic ancestry (n = 400). We used all 1251 individuals in MESA to estimate the proportion of variance in alcohol consumption that was explained by the CpGs, but only used the 588 individuals of EA for the ROC analysis; there were too few heavy drinkers of AA or Hispanic ancestry for meaningful analysis.

The R statistical software (https://www.r-project.org/) was used for all analyses. Linear regression was performed using the function 'lm' for unrelated samples and 'lme' for family samples to account for family structure. LASSO was performed using the function 'glenet' in the 'glenet' R package with the parameter a = 1 and 10-fold cross-validation to select CpGs. The ROC analysis used the 'pROC' package with the 'lme' function for logistic regression, and then the 'predict' function to predict the expected probability, and finally the 'roc' function to estimate sensitivity and specificity of a set of predictors for predicting 'disease' versus 'control' status.

Meta-analysis to identify DNA methylation signature

The inverse variance-weighted random-effects model²⁸ was used in metaanalysis because of the heterogeneity in levels of alcohol consumption and population demographics (Table 1). The meta-analysis was performed in ancestry-stratified whole-blood-derived DNA samples (n=9643 EA and n=2423 AA) and, secondarily, in combined transethnic samples of whole-blood and monocyte-derived DNA (n=13 317). In the meta-analysis, a CpG was further removed if it was missing in five or more studies or its sample size was <20% of the total sample size. We used $P < 0.05/440\ 000 \sim 1 \times 10^{-7}$ to establish significance.

We reported alcohol-related CpGs ($P \le 1 \times 10^{-7}$) in meta-analysis of ancestry-stratified whole-blood-derived DNA samples and in monocytederived DNA samples, and compared alcohol-related CpGs between ancestries and between whole-blood and monocyte samples. We also investigated the DNA methylation levels in several genes that were previously reported to be associated with alcohol metabolism^{29,30} or alcohol-related neurotransmission.³¹⁻³⁴

DNA methylation in former and never drinkers

To investigate if DNA methylation signals differ between 'former' and 'never' drinkers, we performed three additional EWAS analyses with DNA methylation as the outcome variable and three binary traits as the independent variables (adjusting for age, sex, BMI, batch effects and white blood cell counts) in the FHS data. The first analysis using the binary trait 'never' versus 'former' as the independent variable, and the other two the binary traits 'heavy' versus 'never' or 'heavy' versus 'former' as the independent variable. A linear mixed-effects model was used to account for family structure.

Methylation quantitative trait loci analysis. Methylation quantitative trait locus analysis (meQTLs) was performed in three cohorts: FHS (n = 2024), KORA F4 (n = 1799) and the Prospective Investigation of the Vasculature in Uppsala Seniors (PIVUS) study (n = 920). Genotyping, genotype imputation and QC details are described in the Supplementary Information. Using data imputed (allele dosage) to the 1000 Genomes (reference), we selected cis-SNPs (defined as ± 100 kb) with imputation guality score > 0.8 and minor allele frequency ≥ 0.05. The meQTL mapping was performed between the significant alcohol-related CpGs (outcomes) and cis-SNPs (predictors), adjusting for age, sex, BMI, batch effects and white blood cell counts. The proportion of variance (r^2) that can be explained by cis-SNPs or meQTLs for a CpG was also calculated in association analysis. We used a linear model in unrelated individuals or a linear mixed-effects model in family samples to account for familial correlation in association test between an SNP dosage and DNA methylation. Meta-analysis used the inverse-variance weighted random-effects model. We used P < 0.05/n to establish significance, where n was the number of CpG–SNP pairs tested.

Association analysis between methylation and gene expression Gene expression profiling and QC in FHS (n = 1924) and KORA F4 (n = 707) are detailed in the Supplementary Information. To perform the association analysis, the FHS samples were divided into discovery (n = 966) and

replication (n = 958) sets by independent pedigrees. In both FHS and KORA F4 samples, residuals of gene expression levels (Resid_{Gene}) or CpG β -values (Resid_{CpG}) were obtained by adjusting for age, sex, BMI, batch effects and white blood cell counts. Here batch effects and cell proportion differentials (if calculated) were expression-specific or methylation-specific values. The association analysis was then performed between Resid_{Gene} and Resid_{CpG}. A linear model was used in unrelated samples and a linear mixed model was used in family data to account for family structure. The proportion of variance in a transcript that was explained by a CpG was also calculated. Because FHS and KORA F4 used different expression arrays, we only used CpG–gene name pairs that could be matched between the two studies. Therefore, we used the *Z*-score method³⁵ for meta-analysis. We used P < 0.05/n to establish statistical significance, where *n* was the number of CpG–qene transcript pairs tested.

Functional inference and pathway analysis

Genomic features of the alcohol-related CpGs. The genomic location of a CpG provides functional insight into regulatory features.³⁶ According to the annotation 'HumanMethylation450_15017482_v.1.2.csv' provided by Illumina, we compared the enrichment or depletion of several genomic features, including CpG islands, CpG shores and shelves, enhancers, DNA hypersensitivity sites and promoters in the set of alcohol-related CpG probes assessed on the microarray that passed QC. The difference in proportions of a genomic feature was compared by the Fisher's two-sided test.

Gene ontology enrichment analysis and functional inference. We performed Gene Ontology (http://geneontology.org/page/go-enrichmentanalysis) enrichment analysis for the genes that were annotated to the significant alcohol-related CpGs. We also examined the genes whose expression levels were significantly associated with the significant alcoholrelated CpGs.

RESULTS

A methylation biomarker of alcohol consumption

The meta-analysis of the discovery set that included the wholeblood-derived DNA of individuals of EA from eight cohorts (n = 6926; Table 1) identified 361 CpGs at $P < 5 - 10^{-6}$. Of these 361 CpGs, 333 are on the new Infinium MethylationEPIC BeadChip. Using the FHS cohort as the training set, we selected 5 (s = 0.12), 23 (s = 0.08), 78 (s = 'lambda.1se') and 144 (s = 'lambda.min') CpGs out of the 333 CpGs with the LASSO regression (see Materials and Methods). All CpGs in the smaller lists are subsets of the largest set of 144 CpGs (s = 'lambda.min') (Supplementary Table 1). All selected CpGs were available in MESA and ARIC. Five CpGs in the 144 set and one in the 78 CpG set were unavailable in KORA F4 and LBC1936 (Supplementary Table 1).

The most parsimonious set of 5 CpGs explained a substantial proportion of interindividual variance in alcohol consumption in KORA F4 (6.4%), LBC1936 (10.4%), ARIC (5.2%), MESA (9.9%) and FHS (15.0%). The addition of more CpGs yielded larger proportions of explained variance in alcohol consumption. The largest set (144 CpGs) explained 13.1 (KORA F4), 12.0 (LBC1936), 13.8 (ARIC), 13.1 (MESA) and 27.3% (FHS) of variance in alcohol consumption (Table 2). Because the FHS was used as the training cohort to select CpGs, the estimated variance values obtained in the FHS were more optimistic compared with those obtained in the four replication cohorts.

In ROC analysis of 'disease' versus 'control' status (see Materials and Methods), including any CpGs in addition to clinical variables (age, sex and BMI) (the 'Full' model) resulted in a larger AUC value compared with the model with only clinical variables (the 'Null' model). The models with the two smaller sets of CpGs (5 CpGs and 23 CpGs) yielded good prediction (AUC_{Full} > 0.80) in all five cohorts for discriminating heavy drinkers versus non-drinkers; the models with the two larger sets of CpGs (78 CpGs and 144 CpGs) gave good prediction (AUC > 0.80) in all five cohorts for discriminating heavy drinkers versus non-drinkers/light drinkers/ at-risk drinkers, or in discriminating at-risk drinkers versus

Table 2.	The proportion of variance in alcohol consumption
explaine	ed by DNA methylation

Study	Variance explained (%)						
	Null	5 CpGs	23 CpGs	78 CpGs	144 CpGs		
Kora F4 LBC1936 Aric	12.5 9.9 20.0	6.4 10.4 5.2	7.4 11.1 6.0	11.4 12.2 12.4	13.1 12.0 13.8		
MESA FHS	20.0 11.6 7.8	9.9 15.0	10.5 18.9	11.7 24.6	13.0 13.1 27.3		

Abbreviations: ARIC, Atherosclerosis Risk in Communities Study: BMI, body mass index; CpG, cytosine-phosphate-guanine dinucleotide; FHS, Framingham Heart Study; KORA F4, Cooperative Health Research in the Region of Augsburg; LASSO, least absolute shrinkage and selection operator; LBC1936, The Lothian Birth Cohort 1936; MESA, Multi-Ethnic Study of Atherosclerosis. The meta-analysis using the whole-blood-derived DNA of individuals of European ancestry from eight discovery cohorts (n = 6926, see Materials and methods) excluding KORA F4 and LBC1936 identified 361 CpGs with $P < 5 \times 10^{-6}$. Of these 361 CpGs, 333 are on the new Infinium MethylationEPIC BeadChip. Using the FHS data as the training set, we selected 5 (s = 0.12), 23 (s = 0.08), 78 (s = 'lambda.1se') and 144 (s = 'lambda.min') CpGs with the LASSO regression (see Materials and Methods). The testing cohorts included two cohorts of European ancestry with the whole-blood-derived DNA samples (KORA F4 and LBC1936), the African ancestry cohort with the whole-blood-derived DNA samples (ARIC) and the cohort of monocyte-derived DNA samples (MESA) of mixed ancestry (see Table 1). We estimated the proportion of variance in alcohol consumption explained by a list of CpGs as the difference of adjusted R^2 . using a linear regression model that included age, sex and BMI (the 'Null' model) and a model that included a list of CpGs in addition to age, sex and BMI. All selected CpGs were available in MESA and ARIC. Five CpGs in the 144 set and one CpG in the 78 set were unavailable in KORA F4 and LBC1936 (see Supplementary Table 1). The estimated variance values using the FHS data are more optimistic compared with those obtained for the four replication cohorts.

non-drinkers (Figure 2 and Supplementary Figure 1). For example, the addition of the 144 CpGs to the null model yielded a high AUC for discriminating heavy drinkers versus non-drinkers (AUC_{Full} = 0.90–0.99 compared with $AUC_{Null} = 0.63-0.80$) and heavy drinkers versus light drinkers (AUC_{Full} = 0.85-0.99 compared to AUC_{Null} = 0.53-0.61) across the five cohorts; the addition of 78 CpGs to the null model yielded slightly lower AUC values compared with addition of the 144 CpGs: $AUC_{Full} = 0.88-0.99$ in discriminating heavy drinkers versus non-drinkers and AUC_{Full} = 0.82-0.96 in discriminating heavy drinkers versus light drinkers. It is worth noting that in discriminating heavy drinkers versus non-drinkers/ light drinkers, the performance of the 144 CpGs and 78 CpGs was better in MESA and LBC1936 compared with that in FHS (the training cohort); but the performance of these two sets of CpGs was lower in KORA F4 and ARIC (Figure 2). Unavailability of a few CpGs in LBC1936 did not seem to affect discrimination (Table 2 and Figure 2).

Current smoking explained a very small proportion of variance in alcohol consumption. For example, the change in adjusted $R^2 = 0.003$ in FHS and 0.01 in MESA when current smoking was included in the model in addition to age, sex and BMI. Similarly, ALT or AST explained a small proportion of variance in alcohol consumption in FHS: the change in adjusted $R^2 = 0.004$ when either ALS or AST was added to the null model. Therefore, neither ALT nor AST was a good biomarker for alcohol consumption, which was confirmed in ROC analysis: AUC_{Null+ALT or Null+AST} = 0.67 when ALT or AST was added in the null model (AUC_{Null} = 0.66) in discriminating heavy drinkers versus non-drinkers in FHS.

427

Association between DNA methylation and alcohol intake C Liu *et al*

428

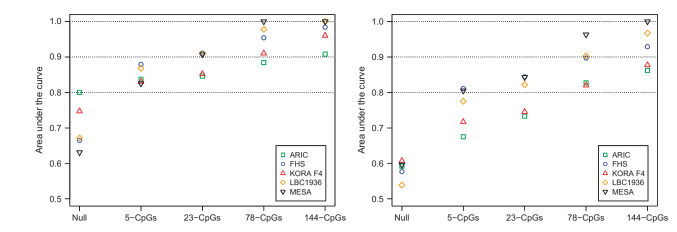


Figure 2. A biomarker of heavy alcohol drinking. Four sets of cytosine-phosphate-guanine dinucleotides (CpGs) were selected at s = 0.12 (5 CpGs), s = 0.08 (23 CpGs), s = 'lambda.1se' (78 CpGs) and s = 'lambda.min' (144 CpGs) using least absolute shrinkage and selection operator (LASSO) in the Framingham Heart Study (FHS) cohort (the training cohort). ROC analysis was performed to classify heavy drinkers versus non-drinkers (left figure) and heavy drinkers versus light drinkers (right figure). 'Non-drinkers' were subjects with no alcohol consumption (i.e., g per day = 0); 'light drinkers' were subjects who consumed 0 < g per day ≤ 28 in men and 0 < g per day ≤ 14 in women; 'heavy drinkers' were subjects who consumed $a \geq 28$ g per day in women. ARIC, The Atherosclerosis Risk in Communities study; KORA F4, The Cooperative Health Research in the Region of Augsburg study; LBC1936, The Lothian Birth Cohort 1936; MESA, The Multi-Ethnic Study of Atherosclerosis.

IlmnID U	CSC gene	Chr	Position	P-value	β	S.e.	UCSC CpG islands	Relation to UCSC CpG island	Enhance
cg03523740 T>	XLNA	1	32 645 027	4.4E – 15	-0.00022	2.8E – 05	Chr 1:32 645 154–32 645 814	N_Shore	
cg20970369 Di	ENND2D	1	111 744 108	3.2E – 12	-0.00023	3.3E – 05	Chr 1:111 746 337-111 747 303	N_Shelf	
cg16246545 PH	HGDH	1	120 255 941	1.5E – 12	-0.00061	8.6E – 05	Chr 1:120 254 844-120 255 499	S_Shore	
cg19266329		1	145 456 128	1.7E – 13	-0.00028	3.8E – 05			TRUE
cg19238380 LA	MNA	1	156 093 948	2.2E – 12	- 0.00029	4.2E – 05			TRUE
cg11194994 PE	EA15	1	160 175 974	7.3E – 15	-0.00017	2.2E – 05	Chr 1:160 175 132-160 175 702	S_Shore	
cg07502661		2	43 398 339	2.6E – 12	-0.00019	2.7E – 05	Chr 2:43 398 040-43398276	S_Shore	
cg00883689 SF	PTBN1	2	54 802 904	3.2E – 12	-0.00028	4.1E-05			TRUE
cg13729116 LE	ETM1	4	1 859 262	6.7E – 18	-0.00018	2.1E-05	Chr 4:1 857 065–1 858 887	S_Shore	
cg25518868 Di	IAPH1	5	140 984 057	2.3E – 12	-0.00012	1.8E – 05			TRUE
cg05593667		6	35 490 744	4.4E – 16	-0.00025	3.1E-05			
cg20732076 TH	RERF1	6	42 335 231	1.5E – 12	-0.00015	2.1E-05			TRUE
cg06189038 G/	AL3ST4	7	99 767 134	4.6E – 13	-0.00016	2.2E – 05	Chr 7:99 768 884–99 769 559	N_Shore	
cg12873476		8	142 402 728	2.8E – 12	-0.00023	3.3E – 05	Chr 8:142 401 533 – 142 402 494	S_Shore	TRUE
cg03599037 Ci	10orf58	10	82 172 508	4.5E – 13	-0.00014	1.9E – 05	Chr 10:82 168 064-82 168 917	S_Shelf	
cg06603309 K0	CNQ1	11	2 724 144	2.7E – 14	0.00017	2.2E – 05	Chr 11:2 720 410–2 722 087	S_Shelf	TRUE
cg11376147 SL	LC43A1	11	57 261 198	9.8E – 13	-0.00026	3.6E – 05			TRUE
cg00271311 Cl	NTF	11	58 389 290	1.6E – 13	-0.00022	2.9E – 05			
cg09448652 SN	NORD30	11	62 621 367	1.3E – 12	-0.00026	3.6E – 05	Chr 11:62 623 359-62 623 877	N_Shore	
cg09737197 Cl	PT1A	11	68 607 675	5.0E – 13	-0.00016	2.2E – 05	Chr 11:68 608 155-68609419	N_Shore	
cg02583484 Hi	NRNPA1	12	54 677 008	1.6E – 19	- 0.00039	4.4E – 05	Chr 12:54 673 322-54 673 550	S_Shelf	
cg23654112 TE	BC1D24	16	2525 928	3.0E – 13	-0.00014	1.9E – 05	Chr 16:2 521 086–2 525 929	Island	
cg08916477 SE	EPT1	16	30 391 350	4.0E – 13	-0.00016	2.2E – 05	Chr 16:30 389 035–30 390 631	S_Shore	
cg06469895 TE	ERF2	16	69 418 206	1.5E – 13	-0.00020	2.8E – 05	Chr 16:69 419 316-69 420 086	N_Shore	
cg00574412 Al	BHD15	17	27 892 866	1.1E – 12	-0.00017	2.3E – 05	Chr 17:27 893 086-27 896 078	N_Shore	
cg21626848 SC	C65	17	39 969 267	3.1E – 15	-0.00023	2.9E – 05	Chr 17:39 967 407-39 968 604	S_Shore	
cg08677210 M	1SI2	17	55 550 613	3.3E – 12	-0.00013	1.9E – 05			TRUE
cg15253293		17	79 366 853	1.1E – 15	-0.00014	1.7E – 05	Chr 17:79 366 806-79374742	Island	
cg24217948 SE	ETBP1	18	42 261 980	2.1E – 12	-0.00028	3.9E – 05	Chr 18:42 258 983–42 260 795	S_Shore	TRUE
cq13127741 C	OMMD7	20	31 331 821	3.1E – 12	-0.00023	3.3E – 05	Chr 20:31 330 957–31 331 410	S Shore	

Abbreviation: CpG, cytosine-phosphate-guanine dinucleotide; S.e, standard error. Epigenome-wide association and meta-analysis of the continuous alcohol intake was performed using all whole-blood-derived DNA samples of European ancestry. The DNA methylation proportion was the outcome variable, grams alcohol consumed per day (g per day) was the predictor variable, adjusting for age, sex, body mass index, technical covariates and white blood cell counts. The inverse-variance weighted random-effects model was performed in meta-analysis (See Supplementary Table 2 for a full set of significant CpGs). The annotation "HumanMethylation450_15017482_v.1.2.csv" provided by Illumina was used to annotate the CpG loci.

Epigenome-wide methylation signature of alcohol intake In the main text, we reported ancestry-stratified meta-analysis $(P < 1 \times 10^{-7})$ for whole-blood-derived DNA in individuals of EA (n = 9643) and AA (n = 2423) (Table 1) using an inverse-variance weighted random-effects model (Figure 1). Meta-analysis of pooled samples $(n = 13 \ 317)$ and several sensitivity tests including

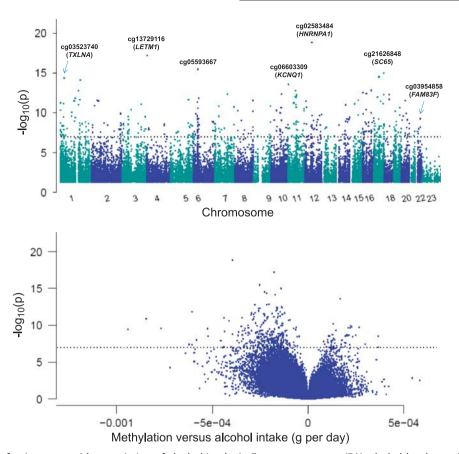


Figure 3. Meta-analysis of epigenome-wide association of alcohol intake in European ancestry (EA) whole-blood samples: the Manhattan plot (top) and the volcano plot (bottom). The DNA methylation proportion was the outcome variable, grams alcohol consumed per day (g per day) was the predictor variable, adjusting for age, sex, body mass index, technical covariates and white blood cell counts. The inverse-variance weighted random-effects model was performed in meta-analysis using all whole blood DNA samples of EA.

EWAS in only drinkers and the investigation of whether or not prevalent CVD or cancer confound the relationship between DNA methylation and alcohol consumption are included in the Supplementary Information.

Genome-wide pattern of DNA methylation associated with alcohol consumption. We identified hundreds of CpGs ($P < 1 \times 10^{-7}$) whose differential methylation across the genome was associated with alcohol intake: 363 CpGs in whole-blood samples of individuals of EA (Table 3, Figure 3 and Supplementary Table 2), CpGs in whole-blood samples of individuals of 165 (AA (Supplementary Table 3 and Supplementary Figure 2) and 62 CpGs in monocyte-derived DNA samples (Supplementary Table 4 and Supplementary Figure 3). Additional CpGs at $P < 1 \times 10^{-1}$ are reported in Supplementary Tables 5-7. Genomic inflation in meta-analysis was estimated at ~10% or less, indicating low additional risk of false-positive findings (Supplementary Table 8). The majority of the alcohol-related CpGs exhibited an inverse relationship between higher alcohol intake and lower methylation (Supplementary Tables 2-4).

Fewer alcohol-related CpGs ($P < 1 \times 10^{-7}$) were identified in the analysis of the categorical alcohol trait that compared light drinkers, at-risk drinkers and heavy drinkers with non-drinkers (Supplementary Tables 9–11 and Supplementary Figures 4–6). Additional CpGs at $P < 1 \times 10^{-4}$ are included in Supplementary Tables 12–14. The majority of the alcohol-related CpGs identified in the analysis of the categorical alcohol trait (Supplementary Tables 9–11) were also significant or nominally significant in the association with the continuous alcohol consumption trait (Supplementary Tables 2–4).

Transethnic replication of methylation signatures

Of the 363 alcohol-related CpGs in EA samples, 56 had P < 0.00014 (0.05/363) in AA samples; of the 165 alcohol-related CpGs in AA samples, 59 had P < 0.00030 (0.05/165) in EA samples. Effect estimates of the 518 (union of 363 and 165) unique CpGs were moderately correlated between EA and AA whole-blood samples: Pearson's correlation r = 0.64 (Figure 4a). For example, cg11376147 in solute carrier family 43 (*SLC43A1*) displayed $P < 1 \times 10^{-7}$ in both EA and AA samples from whole blood (Figure 4b).

Methylation signature in whole-blood- and monocyte-derived DNA

Of the 363 alcohol-related CpGs in EA whole blood samples, 57 replicated (P < 0.00014; 0.05/363) in monocyte samples. Of the 62 alcohol-related CpGs in monocytes, 13 replicated (P < 0.0008; 0.05/62) in whole-blood EA samples. The Pearson's correlation was 0.72 for the 417 unique (union of 363 and 62) CpGs between EA whole-blood samples and monocyte samples (Figure 4c). For example, cg11376147 in *SLC43A1* also displayed $P < 1 \times 10^{-7}$ for association with alcohol consumption in monocyte-derived DNA (Figure 4b).

Similar DNA methylation pattern in former and never drinkers Based on alcohol consumption at prior examinations in FHS, we classified the 693 non-drinkers (Table 1) into 'never' (n = 107) and 'former' drinkers (n = 586). Furthermore, among the 586 'former' drinkers, 91 were 'former' heavy drinkers, 66 were 'former' at-risk drinkers and 429 were 'former' light drinkers. The EWAS using the binary trait 'never' versus 'former' as the independent variable did

Association between DNA methylation and alcohol intake C Liu *et al*

430

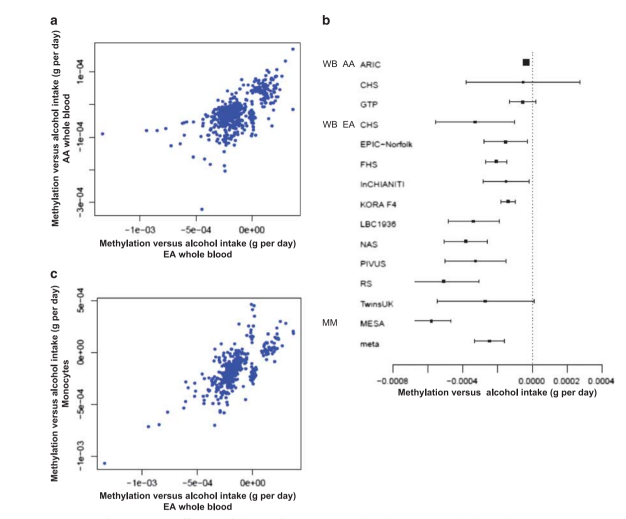


Figure 4. Comparison of regression coefficients of the significant cytosine-phosphate-guanine dinucleotides (CpGs) in association analysis of the continuous alcohol trait (g per day): (a) between European and African whole-blood samples; (b) the Forest plot of effect estimates and standard errors of cg11376147 in all study cohorts; and (c) between European whole-blood and CD14+ monocyte samples. (a) Includes a list of CpGs with $P < 1 \times 10^{-7}$ in EA or AA whole-blood samples and (c) includes a list of CpGs with $P < 1 \times 10^{-7}$ in EA whole-blood samples or in monocyte samples of mixed ancestries. The Pearson's correlation was r = 0.64 between the effect estimates in (a) and r = 0.72 in (c). MM, monocyte, mixed ancestries; WB AA, whole blood, African ancestry; WB EA, whole blood, European ancestry.

not yield any significant results (Supplementary Figure 7). We compared the EWAS results between 'heavy' versus 'never' and 'heavy' versus 'former'. For genome-wide methylation loci, the correlation was 0.32 for regression coefficients and 0.20 for $-\log_{10}$ (*P*-values); for loci with *P*-value $< 1 \times 10^{-7}$ (*n*=92) in either 'heavy' versus 'never' or 'heavy' versus 'former' drinkers, the correlation was 0.91 for regression coefficients and 0.88 for $-\log_{10}$ (*P*-values) (Supplementary Figure 8). These results indicate that DNA methylation levels were not considerably different between 'never' drinkers and 'former' drinkers and that DNA methylation changes due to heavy alcohol consumption revert after individuals abstained from alcohol intake for several years (FHS examinations were ~4 years apart).

Evaluation of smoking in the association between alcohol intake and DNA methylation

It is unclear if current cigarette smoking confounds the association between DNA methylation and alcohol intake, or if smoking and alcohol intake are associated with common CpGs. Therefore, we performed an analysis using smoking as an additional covariate in the EWAS (see Materials and methods). We found that some alcohol-related CpGs displayed a large change (>10% change) in the size of their regression coefficients when smoking was included as an additional covariate in the analysis of wholeblood-derived DNA samples in individuals of EA (35 of the 363 CpGs at $P < 1 \times 10^{-7}$) and AA (92 of the 165 CpGs at $P < 1 \times 10^{-7}$) ancestries, but none of the CpGs in the monocyte-derived DNA samples changed appreciably after additionally adjusting for smoking (Supplementary Tables 2–4). Several of the identified CpGs that displayed a large change in effect estimates following adjustment for smoking, including the CpGs in the aryl-hydrocarbon receptor repressor^{37,38} (Supplementary Tables 5 and 6). We excluded the 35 CpGs that showed large change in effect estimates after adjusting for smoking (Supplementary Table 2) in subsequent analyses that were performed using the whole-blood-derived DNA samples from individuals of EA ancestry.

Alcohol metabolism enzymes and alcohol-related DNA methylation. Several functional DNA sequence variants in the alcohol dehydrogenase (*ADH*) and aldehyde dehydrogenase (*ALDH*) family of genes are known for their effects on alcohol metabolism.^{29,30} We checked CpGs in the introns, exons and regulatory regions in these gene families according to the annotation provided by Illumina. No CpGs in the *ADH* (30 CpGs in seven genes) or *ALDH*

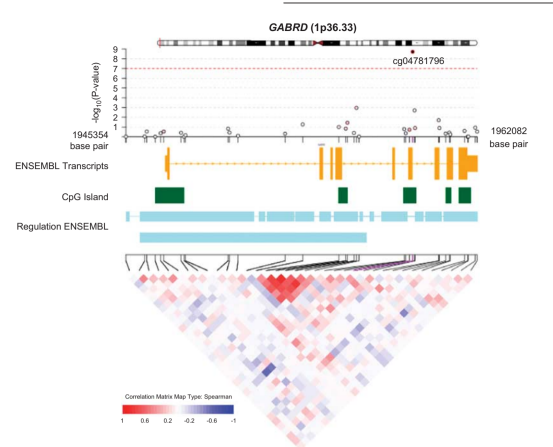


Figure 5. The γ -aminobutyric acid-A (GABA-A) receptor, delta (GABRD): the associations of the 36 cytosine-phosphate-guanine dinucleotides (CpGs) within GABRD, genomic and regulatory features and correlation of methylation measurements. The results were obtained in metaanalysis of the association analysis of 9643 whole-blood-derived DNA samples of European ancestry (EA) individuals. The correlation of these 36 CpGs was calculated using the methylation measurements at 36 CpGs, adjusting for age, sex, technical covariates and white cell blood counts in the Framingham Heart Study samples.

(340 CpGs in 19 genes) loci were differentially methylated $(P < 1 \times 10^{-7})$ in relation to alcohol use (Supplementary Tables 5-7, 15, 16).

Neurotransmitter receptors and alcohol-related DNA methylation y-aminobutyric acid (GABA), a major neurotransmitter, and its receptors are known for their involvement in the acute and chronic behavioral effects of ethanol in humans and animal models.^{31–34} A total of 607 CpGs were mapped to introns, exons and regulatory regions of 22 GABA receptor genes (Supplementary Table 17). In meta-analysis of whole-blood-derived DNA samples of EA individuals, cg04781796 ($\beta_{alcohol} = 0.0002$, $P = 1.9 \times 10^{-9}$) and cg09577455 ($\beta_{alcohol} = -0.0015$, $P = 3.0 \times 10^{-8}$) were significantly associated with alcohol consumption (Supplementary Table 2). However, neither of these CpGs was significant in whole-bloodderived DNA samples of people of AA (cg04781796: $\beta_{\text{alcohol}} = 0.00004, P = 0.17;$ cg09577455: $\beta_{\text{alcohol}} = -0.00003,$ P = 0.0016) or in monocyte-derived DNA samples (cg04781796: $\beta_{\text{alcohol}} = 0.0001$, P = 0.22; cg09577455: $\beta_{\text{alcohol}} = -0.00002$, P = 0.82). The CpG cg04781796 is located in a CpG island (Chr 1: 1 959 414-1 959 867, hg19) that is intronic to the GABA-A receptor, delta (GABRD; Figure 5); and cg09577455 is located in the north shore of a CpG island (Chr 6: 29 595 298-29 595 795, hq19) that is intronic to GABA-B receptor subunit 1 (GABBR1, Chr 6; Supplementary Figure 9). CpG cg04781796 displayed higher methylation in relation to greater alcohol intake, whereas cq09577455 displayed lower methylation levels in association with increased alcohol intake.

Genetic basis underlying the significant alcohol-related CpGs We tested for association of the methylation levels of 328 CpGs (selected in meta-analysis of DNA of EA individuals in Supplementary Table 2) with nearby SNPs (±100KB, cis-SNPs or cis-meQTLs) in three cohorts (total number of individuals = 4623 from FHS, KORA F4 and PIVUS) (see Materials and methods). A total of 105 759 SNP-CpG pairs were formed from these 328 CpGs with cis-SNPs. Meta-analysis of the FHS, KORA F4 and PIVUS samples identified 14 160 SNP-CpG pairs (170 unique CpGs and 12 857 unique *cis*-SNPs) at $P < 0.05/105759 \sim 4.7 \times 10^{-100}$ (Supplementary Table 18). We found that 16 CpGs had meQTLs that explained 20-61% of interindividual variance in methylation at the corresponding CpG (Supplementary Table 18). None of these 12 857 significant meQTLs was associated with alcoholrelated traits ($P < 1 \times 10^{-8}$) by querying these significant meQTLs in the Catalog of Published Genome-Wide Association Studies (http://www.genome.gov/gwastudies/, latest version released on May 12, 2015).

Association of alcohol-related DNA methylation with gene expression

We tested for associations between the 328 alcohol-related CpGs and blood gene expression levels in FHS (n = 1924) and KORA F4 (n = 707) for genes within 1 Mb of these 328 CpGs in both studies (see Materials and methods). Meta-analysis identified 110 CpGgene pairs (83 unique CpGs and 100 unique genes) at P < 0.05- $77111 = 7 \times 10^{-6}$, where 7111 is the number of tested CpG-gene pairs (Supplementary Table 19). Of the 110 significant pairs, 86

431

(78%) displayed negative correlations between methylation and mRNA levels.

We examined associations of the two significant alcohol-related CpGs in GABA receptor genes with expression of *cis* genes. At the GABRD locus, cq04781796 was not associated with expression of any genes in blood within 1 Mb, whereas cg09577455 in the GABBR1 locus was associated with the expression of the interferon-induced transmembrane protein 4 pseudogene (IFITM4P; $P = 2.4 \times 10^{-6}$) (Supplementary Table 19). Owing to the important role of GABA receptors in alcohol-induced signal transduction and immune functions, we carried out additional association analyses between these two CpGs and gene transcripts beyond 1 Mb or on different chromosomes (i.e. trans associations) in both FHS and KORA F4. Of the 35 746 association significant pairs. 228 showed association (P < 0.05 - $/35746 \sim 1.4 \times 10^{-6}$) with methylation of cg04781796 (Supplementary Table 20) and 13 transcripts were associated with cg09577455 (Supplementary Table 21) in the meta-analysis.

Functional inference and pathway analysis

Genomic features of the alcohol-related CpGs. We found that the 328 alcohol-related CpG set was significantly enriched for CpG island shores (48% versus 24%, $P = 7.3 \times 10^{-12}$) and enhancers (29% versus 22%, P = 0.003) compared with all CpGs that passed QC in meta-analysis. In contrast, the 328 alcohol-associated CpG set was significantly depleted for CpG islands (16% versus 32%, $P = 1.1 \times 10^{-6}$) and promoters (3% versus 7%; P = 0.009) (Supplementary Table 22). There was no significant difference in proportions of CpG island shelves and DNase I hypersensitive sites among the 328 alcohol-associated CpGs.

We found similar enrichment and depletion for the 144 CpGs that were selected in biomarker analysis. These 144 CpGs were significantly enriched for CpG island shores (47% versus 24%, $P = 4.5 \times 10^{-11}$) and enhancers (30% versus 22%, P = 0.002), but significantly depleted for CpG islands (13% versus 32%, $P = 1.0 \times 10^{-6}$) and promoters (2% versus 7%, P = 0.007).

Gene ontology enrichment analysis and functional inference.

A total of 257 genes were annotated to the 328 alcohol-related CpGs. These 257 genes were enriched for 95 biological processes (Bonferroni-corrected P < 0.05, Supplementary Table 23) including regulation of transcription, macromolecule metabolic process and cellular response to stress and chemicals. The most significant biological process constituted 32 genes enriched (>4-fold) for 'negative regulation of transcription from RNA polymerase II promoter' (Bonferroni-corrected $P = 2.3 \times 10^{-7}$; Supplementary Table 24). For the 100 cis genes whose transcript levels were significantly associated with 83 CpGs (Supplementary Table 19), the analysis of biological processes showed that the most significantly enriched process was 'negative regulation of transposition' (Bonferronicorrected $P = 8.0 \times 10^{-4}$, Supplementary Table 25). Other enriched processes included defense response to virus (P = 0.006) and DNA cytosine deamination (P=0.02). The trans-transcripts that were significantly associated with cg04781796 (GABRD) were enriched for pathways that are involved in immune functions such as lymphocyte activation $(P = 1.1 \times 10^{-11})$ and immune system process $(P=3.2\times10^{-11})$; Supplementary Table 26). The *trans*-transcripts that were associated with cq09577455 in GABBR1 were also enriched for immune response (P = 0.015; Supplementary Table 27).

DISCUSSION

We conducted an EWAS of alcohol intake in 13 cohorts including 13 317 samples of whole-blood or monocyte-derived DNA from individuals of mostly EA and AA. We identified hundreds of differentially methylated CpGs ($P < 1 \times 10^{-7}$) in relation to alcohol consumption. More than half of the alcohol-related methylation

sites were associated with *cis*-genetic variants, supporting the hypothesis that DNA methylation sites are sensitive to both environmental and genetic influences.³⁹ In addition, we developed a robust and replicable DNA methylation biomarker that provides substantial discrimination for current heavy alcohol intake.

A set of 144 CpGs was highly predictive for discriminating current heavy alcohol drinkers from non-drinkers (AUC > 0.90) in all replication cohorts. As a biomarker, these selected CpGs performed better than commonly clinical variables and biomarkers in discriminating current heavy alcohol drinking.⁷ This is in line with the discriminatory power of DNA methylation for other complex traits, such as BMI.⁴⁰ Therefore, a whole-blood DNA methylation biomarker has the potential to be developed into a commercially marketable diagnostic test to detect current heavy alcohol consumption. Such a test could be useful to supplement and validate self-reported alcohol consumption data, or in a forensic setting, or as a screening test.

The biomarker analysis and ancestry-stratified meta-analysis showed that a number of DNA methylation sites displayed consistent alcohol-related effects in whole-blood samples of people of EA and AA. However, the transancestry comparison also showed the lack of similarities of many CpG sites. We propose three explanations. First, some DNA methylation sites are truly ancestry-specific, which needs to be confirmed by future studies. Second, sample heterogeneity in alcohol consumption may explain a part of the non-concordance for some CpGs in AA and EA groups. For example, in ARIC, ~76% individuals were non-drinkers and ~17% were heavy drinkers, whereas in most EA cohorts, >60% of participants were light drinkers. Third, a large difference in sample sizes (EA n = 9643 and AA n = 2423) and the probability in sampling are additional reasons for the lack of replication when a Bonferroni-corrected threshold was used.

We provide evidence that alcohol-related DNA methylation is associated with gene expression in whole blood. Of note, we showed that whole-blood epigenetic changes in GABA receptor genes were significantly associated with the expression levels of a number of genes that are involved in immune function supporting the recent findings that GABA and its receptor have effects on immune cells through cross-talk between the nervous system and the immune system.⁴¹ However, as our data are cross-sectional and observational in nature, further research is needed to determine if these changes are causal or reactive. The gene set analysis is based on a crucial and unrealistic independence assumption pertaining to genes, which may not be valid for biological processes. Therefore, we should interpret the significant *P*-values with caution.⁴²

In addition to the cross-sectional nature of this study, our findings were limited to DNA samples from mostly middle- and older-aged individuals of EA and AAs. Future studies are needed to investigate the generalizability of our findings to other age groups and ancestries. Nevertheless, as the largest study of its kind, this work identified a robust alcohol-related DNA methylation signature in blood and demonstrated that the alcohol-related methylation changes in blood are of sufficient magnitude to be interesting clinically, which addresses a gap within the field. Future studies are warranted to investigate whether alcohol-related methylation in blood affects GABA neurotransmitter function in the brain and to investigate how alcohol-related epigenetic modifications influence the beneficial and detrimental downstream consequences of alcohol-related health outcomes. Identifying how alcohol-induced DNA methylation changes modify gene expression and result in pathway activation or suppression may shed light on the molecular basis of alcohol addiction and alcohol-related diseases and reveal new therapeutic strategies.

CONFLICT OF INTEREST

Erik Ingelsson is an advisor and consultant for Precision Wellness Inc (Redwood City, CA). The remaining authors declare no conflict of interest.

ACKNOWLEDGMENTS

Detailed acknowledgments were included in Supplementary Information.

ACCESSION CODES. The meta-analysis results for the continuous and categorical alcohol consumption traits can be downloaded from the database of Genotypes and Phenotypes (dbGaP) CHARGE Summary site under accession phs000930. DNA methylation measurements by several cohorts have been deposited in dbGap. Contact information for obtaining DNA methylation measurements from the European cohorts and some of the American cohorts is available upon request.

REFERENCES

- 1 NIAAA. Alcohol Facts and Statistics. Available at: https://www.niaaa.nih.gov/ alcohol-health/overview-alcohol-consumption/alcohol-facts-and-statistics.
- 2 Rehm J, Baliunas D, Borges GL, Graham K, Irving H, Kehoe T *et al*. The relation between different dimensions of alcohol consumption and burden of disease: an overview. *Addiction* 2010; **105**: 817–843.
- 3 Ogeil RP, Room R, Matthews S, Lloyd B. Alcohol and burden of disease in Australia: the challenge in assessing consumption. *Aust NZ J Public Health* 2015; **39**: 121–123.
- 4 Rehm J, Taylor B, Roerecke M, Patra J. Alcohol consumption and alcoholattributable burden of disease in Switzerland, 2002. *Int J Public Health* 2007; **52**: 383–392.
- 5 Ferreira-Borges C, Rehm J, Dias S, Babor T, Parry CD. The impact of alcohol consumption on African people in 2012: an analysis of burden of disease. *Trop Med Int Health* 2015; **21**: 52–60.
- 6 Allen JP. Use of biomarkers of heavy drinking in health care practice. *Mil Med* 2003; **168**: 364–367.
- 7 Liangpunsakul S, Lai X, Ross RA, Yu Z, Modlik E, Westerhold C et al. Novel serum biomarkers for detection of excessive alcohol use. Alcohol Clin Exp Res 2015; 39: 556–565.
- 8 Zahs A, Curtis BJ, Waldschmidt TJ, Brown LA, Gauthier TW, Choudhry MA *et al.* Alcohol and epigenetic changes: summary of the 2011 Alcohol and Immunology Research Interest Group (AIRIG) meeting. *Alcohol* 2012; **46**: 783–787.
- 9 Weng JT, Wu LS, Lee CS, Hsu PW, Cheng AT. Integrative epigenetic profiling analysis identifies DNA methylation changes associated with chronic alcohol consumption. *Comput Biol Med* 2015; 64: 299–306.
- 10 Leake I. Liver disease: alcohol causes epigenetic changes in hepatic stellate cells. Nat Rev Gastroenterol Hepatol 2014; 11: 704.
- 11 Nieratschker V, Batra A, Fallgatter AJ. Genetics and epigenetics of alcohol dependence. J Mol Psychiatry 2013; 1: 11.
- 12 Robison AJ, Nestler EJ. Transcriptional and epigenetic mechanisms of addiction. *Nat Rev Neurosci* 2011; **12**: 623–637.
- 13 Robertson KD, Uzvolgyi E, Liang G, Talmadge C, Sumegi J, Gonzales FA et al. The human DNA methyltransferases (DNMTs) 1, 3a and 3b: coordinate mRNA expression in normal tissues and overexpression in tumors. *Nucleic Acids Res* 1999; 27: 2291–2298.
- 14 Harlaar N, Hutchison KE. Alcohol and the methylome: design and analysis considerations for research using human samples. *Drug Alcohol Depend* 2013; 133: 305–316.
- 15 Zhang R, Miao Q, Wang C, Zhao R, Li W, Haile CN et al. Genome-wide DNA methylation analysis in alcohol dependence. Addict Biol 2013; 18: 392–403.
- 16 Clark SL, Aberg KA, Nerella S, Kumar G, McClay JL, Chen W et al. Combined whole methylome and genomewide association study implicates CNTN4 in alcohol use. Alcohol Clin Exp Res 2015; 39: 1396–1405.
- 17 Zhao R, Zhang R, Li W, Liao Y, Tang J, Miao Q *et al.* Genome-wide DNA methylation patterns in discordant sib pairs with alcohol dependence. *Asia Pac Psychiatry* 2013; **5**: 39–50.
- 18 Philibert RA, Plume JM, Gibbons FX, Brody GH, Beach SR. The impact of recent alcohol use on genome wide DNA methylation signatures. *Front Genet* 2012; **3**: 54.
- 19 Chen YA, Lemire M, Choufani S, Butcher DT, Grafodatskaya D, Zanke BW *et al.* Discovery of cross-reactive probes and polymorphic CpGs in the Illumina Infinium HumanMethylation450 microarray. *Epigenetics* 2013; **8**: 203–209.
- 20 Loomba R, Hwang SJ, O'Donnell CJ, Ellison RC, Vasan RS, D'Agostino RB Sr. et al. Parental obesity and offspring serum alanine and aspartate aminotransferase levels: the Framingham Heart Study. *Gastroenterology* 2008; **134**: 953–959.
- 21 Jung M, Pfeifer GP. Aging and DNA methylation. BMC Biol 2015; 13: 7.

- 22 Zhang FF, Cardarelli R, Carroll J, Fulda KG, Kaur M, Gonzalez K et al. Significant differences in global genomic DNA methylation by gender and race/ethnicity in peripheral blood. *Epigenetics* 2011; 6: 623–629.
- 23 Dick KJ, Nelson CP, Tsaprouni L, Sandling JK, Aissi D, Wahl S et al. DNA methylation and body-mass index: a genome-wide analysis. *Lancet* 2014; 383: 1990–1998.
- 24 Bock C. Analysing and interpreting DNA methylation data. *Nat Rev Genet* 2012; **13**: 705–719.
- 25 Houseman EA, Kelsey KT, Wiencke JK, Marsit CJ. Cell-composition effects in the analysis of DNA methylation array data: a mathematical perspective. *BMC Bioinform* 2015; 16: 95.
- 26 Teschendorff AE, Zhuang J, Widschwendter M. Independent surrogate variable analysis to deconvolve confounding factors in large-scale microarray profiling studies. *Bioinformatics* 2011; 27: 1496–1505.
- 27 Price AL, Patterson NJ, Plenge RM, Weinblatt ME, Shadick NA, Reich D. Principal components analysis corrects for stratification in genome-wide association studies. *Nat Genet* 2006; **38**: 904–909.
- 28 Borenstein M, Hedges LV, Higgins JP, Rothstein HR. A basic introduction to fixed-effect and random-effects models for meta-analysis. *Res Synth Methods* 2010; 1: 97–111.
- 29 Zakhari S. Overview: how is alcohol metabolized by the body? Alcohol Res Health 2006; 29: 245–254.
- 30 Takeshita T, Morimoto K, Mao XQ, Hashimoto T, Furuyama J. Phenotypic differences in low Km aldehyde dehydrogenase in Japanese workers. *Lancet* 1993; 341: 837–838.
- 31 Olsen RW, Hanchar HJ, Meera P, Wallner M. GABAA receptor subtypes: the 'one glass of wine' receptors. *Alcohol* 2007; **41**: 201–209.
- 32 Kumar S, Porcu P, Werner DF, Matthews DB, Diaz-Granados JL, Helfand RS *et al.* The role of GABA(A) receptors in the acute and chronic effects of ethanol: a decade of progress. *Psychopharmacology (Berl)* 2009; **205**: 529–564.
- 33 Colombo G, Addolorato G, Agabio R, Carai MA, Pibiri F, Serra S et al. Role of GABA (B) receptor in alcohol dependence: reducing effect of baclofen on alcohol intake and alcohol motivational properties in rats and amelioration of alcohol withdrawal syndrome and alcohol craving in human alcoholics. *Neurotox Res* 2004; 6: 403–414.
- 34 Agabio R, Colombo G. GABAB receptor ligands for the treatment of alcohol use disorder: preclinical and clinical evidence. *Front Neurosci* 2014; **8**: 140.
- 35 Zaykin DV. Optimally weighted Z-test is a powerful method for combining probabilities in meta-analysis. J Evol Biol 2011; 24: 1836–1841.
- 36 Wagner JR, Busche S, Ge B, Kwan T, Pastinen T, Blanchette M. The relationship between DNA methylation, genetic and expression inter-individual variation in untransformed human fibroblasts. *Genome Biol* 2014; 15: R37.
- 37 Joubert BR, Haberg SE, Nilsen RM, Wang X, Vollset SE, Murphy SK et al. 450 K epigenome-wide scan identifies differential DNA methylation in newborns related to maternal smoking during pregnancy. Environ Health Perspect 2012; 120: 1425–1431.
- 38 Philibert RA, Beach SR, Brody GH. Demethylation of the aryl hydrocarbon receptor repressor as a biomarker for nascent smokers. *Epigenetics* 2012; 7: 1331–1338.
- 39 Liu Y, Li X, Aryee MJ, Ekstrom TJ, Padyukov L, Klareskog L et al. GeMes, clusters of DNA methylation under genetic control, can inform genetic and epigenetic analysis of disease. Am J Hum Genet 2014; 94: 485–495.
- 40 Shah S, Bonder MJ, Marioni RE, Zhu Z, McRae AF, Zhernakova A et al. Improving phenotypic prediction by combining genetic and epigenetic associations. *Am J Hum Genet* 2015; **97**: 75–85.
- 41 Jin Z, Mendu SK, Birnir B. GABA is an effective immunomodulatory molecule. *Amino Acids* 2013; 45: 87–94.
- 42 Goeman JJ, Buhlmann P. Analyzing gene expression data in terms of gene sets: methodological issues. *Bioinformatics* 2007; 23: 980–987.

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 4.0 International License. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http:// creativecommons.org/licenses/by-nc-nd/4.0/

© The Author(s) 2018

Supplementary Information accompanies the paper on the Molecular Psychiatry website (http://www.nature.com/mp)