

Supplementary Information

GWAS of risk tolerance and risky behaviors in over one million individuals identify hundreds of loci and reveal shared genetic influences

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1 Study overview and phenotype definitions

1.1 Study motivation

Risk tolerance, or the willingness to take risks to obtain rewards, is a fundamental parameter in economics, finance, and behavioral decision theory. Different measures of risk preferences have previously been linked to real-world behaviors such as portfolio allocation and occupational choice, as well as health behaviors and addiction phenotypes such as smoking, exercising, and alcohol and drug use^{1–5}. Further, recent work has demonstrated the existence of a reliable general factor of risk preference that is generalizable both to specific and real-world risky behaviors⁶.

Elucidating the determinants of individual differences in general risk tolerance is an active field of research^{3,4}. General risk tolerance has been found to be moderately heritable in twin studies ($h^2 \sim 30\%$), although heritability estimates in the literature vary, ranging from 20% to 60%^{2,7–9}; an earlier study based on molecular genetic data had confirmed the heritability of the trait¹⁰. Some previous studies have attempted to identify specific genetic variants that are associated with general risk tolerance. However, most of these attempts have been conducted in relatively small samples with a few hundred to at most a few thousand individuals^{11–14}; see **Supplementary Table 11.1**. Given that the effect of any specific single nucleotide polymorphism (SNP) on a genetically complex trait like general risk tolerance is likely to be extremely small^{10,15–18}, these earlier studies were most likely underpowered. Low statistical power not only implies a low probability to detect true effects, but also a high chance of finding false positives, a strong tendency to overestimate the effects of statistically significant variables, and a high likelihood that significant findings will have the wrong sign¹⁹. Accordingly, the replication record of these underpowered studies has been disappointing^{20,21}.

The purpose of this study is to investigate the molecular genetic architecture of general risk tolerance, adventurousness, and of a number of risky behaviors and to identify specific genetic variants associated with the phenotypes in well-powered genome-wide association studies (GWAS). Our findings could help elucidate the genetic and biological mechanisms that underlie individual variation in the willingness to avoid or engage in risky behavior.

1.2 Phenotype definitions

1.2.1 *General risk tolerance*

Our main phenotype is self-reported “general risk tolerance,” defined as one’s tendency or willingness to take risks in general. For our discovery stage, we combine data from the UK Biobank and from the 23andMe cohort.

We use the following survey question from the UK Biobank ($n = 431,126$):

“Would you describe yourself as someone who takes risks? Yes / No.”

Throughout the study all risk-related phenotypes are coded so that a higher risk tolerance is associated with a higher phenotype value. For example, in the UKB “yes” is coded as 1 and “no” as 0. The majority of the 431,126 respondents in the UKB were only assessed once, while a subset of 18,102 individuals answered the survey question a second time. 15,618 of these re-assessed individuals gave consistent answers in both waves, while 2,484 changed their responses. When the

answer between the two assessments changed, we took the average response (i.e., 0.5) as the individual's measure of risk tolerance. Consistent with prior research^{22,23}, a much higher fraction of males (34%) than females (19%) in the UKB cohort described themselves as risk tolerant on the general-risk-tolerance measure ($P < 1 \times 10^{-100}$, t -test; **Extended Data Fig. 1.1**).

In the 23andMe cohort, our main phenotype is again self-reported “general risk tolerance.” We use the survey question ($n = 508,782$):

“In general, people often face risks when making financial, career, or other life decisions. Overall, do you feel comfortable or uncomfortable taking risks? [1] Very comfortable / [2] Somewhat comfortable / [3] Neither comfortable nor uncomfortable / [4] Somewhat uncomfortable / [5] Very uncomfortable.”

We reverse this coding for our GWAS, so that “1” is coded as the least risk-related response category and “5” as the most risk-related response category.

In total, then, our full general-risk-tolerance discovery meta-analysis includes 939,908 people from the UK Biobank and from the 23andMe cohort.

For our replication stage ($n = 35,445$), we combined 10 independent cohorts from seven studies with survey questions on general risk tolerance, all of which included a question similar to the one asked in the UKB or in Dohmen *et al.*⁴:

“How do you see yourself: are you generally a person who is fully prepared to take risks or do you try to avoid taking risks? Please tick a box on the scale, where the value 0 means: ‘not at all willing to take risks’ and the value 10 means: ‘very willing to take risks’.”

In **Supplementary Table 1.1** we report an overview of the cohorts included in our study. **Supplementary Table 1.2** lists the detailed general-risk-tolerance survey questions available in each cohort. The UKB is the only cohort that asks the general-risk-tolerance question in a binary fashion; the 23andMe cohort asked this question on a 5-point Likert scale, and all seven replication cohorts asked their participants this question on a 10- or 11-point Likert scale. All of the replication cohorts are cross-sectional, and only the UKB includes more than one measurement over time for some individuals.

As we further describe in **Supplementary Information sections 3.3** and **7.4**, we used bivariate LD Score Regression²⁴ to estimate the genetic correlation between: (1) the UK Biobank risk-tolerance GWAS and the 23andMe risk-tolerance GWAS; (2) the UK Biobank risk-tolerance GWAS and the replication GWAS; (3) the 23andMe risk-tolerance GWAS and the replication GWAS; and (4) the full discovery GWAS of general risk tolerance (UK Biobank + 23andMe) and the replication GWAS. For the last three correlations, we see significant, moderately high, and positive genetic correlations between 0.75 and 0.83 that are indistinguishable from unity. However, for (1), we find a genetic correlation that *is* distinguishable from unity ($\hat{r}_g = 0.767$, $SE = 0.021$). Though this genetic correlation is lower than expected, it is high enough to justify the meta-analysis of the UK Biobank and 23andMe summary statistics for general risk tolerance that we perform as our discovery GWAS¹⁸. However, the imperfect correlation points to some degree of heterogeneity across the cohorts, which may attenuate the genetic signals we can observe in our study.

We note that there are various alternative ways to measure risk tolerance, including behavioral experiments with real stakes²⁵, hypothetical choices¹, and survey questions²⁵. The phenotypic

correlation between these measures is typically moderate^{2,4,26}, although correcting for measurement error substantially increases these correlations^{2,26}.

Our approach of using survey measures of general risk tolerance has several advantages. First, these measures have been shown to correlate with a wide range of risky behaviors such as investments in stocks, active sports, self-employment, and smoking, even after controlling for age, sex, education, wealth, and income^{1,2,4,26}. Thus, these survey measures capture an important dimension of individual differences. There is evidence that measures of general risk tolerance are good all-around predictors of risky behavior and perform better in this respect than more specific survey measures of risk tolerance⁴. Second, survey questions of general risk tolerance are cheap and easy to collect and have been added to numerous questionnaires, including the UKB, thus allowing us to reach a large enough sample size to conduct a well-powered GWAS.

Although general risk tolerance may be closely related to some psychological measures of personality such as extraversion, novelty seeking, or sensation seeking, these constructs are not identical. For this reason, personality measures were not included in our main GWAS of general risk tolerance, but below we study the predictive power of a polygenic score of general risk tolerance for several of these psychological measures of personality, as well as the genetic correlation between general risk tolerance and some of these measures.

We also performed supplementary GWAS for adventurousness, four risky behaviors in the UK Biobank (automobile speeding propensity, drinks per week, ever smoker, and number of sexual partners), and the first principal component of these four risky behaviors. Moving forward, we refer to the main GWAS of general risk tolerance as our “primary GWAS” and the GWAS of these additional six phenotypes as our “supplementary GWAS.”

1.2.2 *Adventurousness*

We also performed a GWAS of adventurousness, since this phenotype is known to be related to risk-taking behavior²⁷. For this GWAS, we use only summary statistics from the 23andMe cohort, and we use the following survey question ($n = 557,923$):

“If forced to choose, would you consider yourself to be more cautious or more adventurous? [1] Very cautious / [2] Somewhat cautious / [3] Neither / [4] Somewhat adventurous / [5] Very adventurous.”

We maintain this coding, where “1” is coded as the least risk-related response category and “5” as the most risk-related response category.

1.2.3 *Other supplementary UKB Risky Behaviors*

We also conducted GWAS of four self-reported risky behaviors in the UKB. Our selection strategy for these phenotypes was twofold. As a first step, we chose a set of potential GWAS phenotypes by searching the UKB database for risky behaviors across various domains. The risky behaviors we originally considered included automobile speeding propensity, use of sun protection, age of first sexual intercourse, number of lifetime sexual partners, teenage conception (females only), as well as whether an individual was ever a tobacco smoker, whether an individual is a former tobacco smoker, age of tobacco smoking onset, number of cigarettes per day. We also considered whether an individual was ever an alcohol drinker, whether an individual is a current alcohol drinker, whether an individual is an excessive alcohol drinker, and number of drinks per day. (Ultimately,

we decided not to retain the use of sun protection in our main analyses, because there was no significant genetic correlation between this phenotype and the preliminary results from our GWAS of general risk tolerance using the first release of UKB data ($\hat{r}_g = 0.025$, P value = 0.701, as estimated using bivariate LD Score regression).)

Our decision to consider these phenotypes for inclusion in our study builds on previous studies from various disciplines showing that measures of risk tolerance correlate with a range of risky behaviors across various domains. For example, economists tend to think of risky behaviors such as drinking, smoking, or fast driving as choices associated with uncertain schedules of benefits and costs, and that should thus correlate with risk tolerance. Consistent with this perspective, empirical studies have documented that risk tolerance correlates positively with such risky behaviors^{1–5,28}. However, we highlight that epidemiologists, psychologists and public health researchers typically view smoking and drinking as addiction phenotypes and correlates of mental health^{29,30}. Addictive behaviors such as heavy drinking have been shown to co-occur with a disinhibited personality style (i.e., a personality style associated with behavioral disinhibition, antisocial behavior, and sexual promiscuity), conduct disorder (e.g., rule breaking, criminality, and reckless driving), and attention deficit hyperactivity disorder (ADHD). The co-occurrence of these traits seems to be partly due to a highly heritable latent factor that psychologists call “externalizing”^{31–34}. Thus, the drinking and smoking phenotypes we considered are likely to capture both normal-range variation in risk tolerance among people and addictive or externalizing behaviors. Because we are primarily interested in drinking and smoking due to their close relationship to risk tolerance, we refer to these phenotypes as “risky behaviors” in most instances in the main text and in the rest of this **Supplementary Information**^a. Nonetheless, when interpreting our results it should be kept in mind that both drinking and smoking are also addiction or externalizing phenotypes (for example, the positive genetic correlation we find between risk tolerance and drinks per week could conceivably reflect a correlation between risk tolerance and the addiction or externalizing component, rather than the risk tolerance component, of drinks per week).

As a second step in our phenotype selection process, we selected just one phenotype in each domain of risky behavior (i.e., driving behavior, drinking behavior, smoking behavior, and sexual behavior). To do so, we prioritized: 1) phenotypes available in the entire UKB sample, since our general-risk-tolerance phenotype is defined for everyone in the UKB sample; 2) phenotypes that had been previously explored in other published GWAS; and 3) phenotypes which showed a high phenotypic correlation with our main general-risk-tolerance phenotype in the first release of the UKB data (this is the data we had access to when deciding which phenotypes to select). If not reported in the text below, these correlations from the first release of the UKB are available upon request, although the differences in the correlations between the first and full release are small. For a list of the phenotypic correlations between the selected phenotypes in the full release of the UKB, see **Supplementary Table 1.3**. Below we highlight how the phenotypes in each domain of risky behavior were selected and are defined.

Automobile speeding propensity: Automobile speeding propensity is the only phenotype available that measures risky driving behavior in the UKB; its phenotypic correlation with general risk tolerance in the first release of the UKB is 0.164. Respondents were asked, “How often do you drive faster than the speed limit on the motorway?” Response options include: 1) Never/rarely; 2)

^a In the case of smoking, as we discuss below, the *ever smoker* measure we analyze captures smoking initiation (rather than smoking intensity), which may be particularly closely related to risk tolerance given the associated health risks.

Sometimes; 3) Often; 4) Most of the time; and 5) Do not drive on the motorway. We first dropped all participants who reported not driving on the motorway, and then we normalized^a our categorical variable for males and females separately. In total, this GWAS includes 404,291 individuals in the UK Biobank.

Drinks per week: There are several phenotype options that measure drinking behavior in the UKB. After considering only phenotypes that cover the entire UKB sample, we were left with two: drinks per week and excessive alcohol drinking. Our drinks per week measure is constructed from responses to a sequence of questions. First, respondents were asked how often they drink alcohol, and response options include 1) daily or almost daily; 2) three or four times per week; 3) once or twice per week; 4) one to three times per month; 5) special occasions only; and 6) never. Respondents who reported drinking once per week or more were asked how many glasses of various types of alcoholic beverages they consume per week. We used the sum of all alcoholic drinks per week as our drinks per week phenotype for these respondents. Respondents who reported drinking less than once per week (one to three times per month or on special occasions only) were asked how many glasses of various types of alcoholic beverages they consume per *month*. For these respondents, we added the total number of drinks per month and divided by 4 to arrive at an approximated number of drinks per week. Respondents who reported never drinking were coded as 0.

For the excessive alcohol drinking phenotype, we use current UK Chief Medical Officer drinking guidelines^b to code respondents who drink 14 or fewer drinks per week as 0 and more than 14 drinks per week as 1. Our drinks per week phenotype had a higher phenotypic correlation with general risk tolerance in the first release of the UKB (0.139) than did our excessive alcohol drinking phenotype (0.074); further, drinks per week is a consistent phenotype studied in the alcohol GWAS literature. We therefore use this drinks per week phenotype as our final drinking behavior GWAS phenotype. In total, this GWAS includes 414,343 individuals in the UK Biobank.

Ever smoker: There are several potential measures of smoking behavior in the UKB. These include: 1) ever-tobacco smoker status; 2) former tobacco smoker status (among ever-tobacco smokers); 3) age of tobacco smoking onset (among ever-tobacco smokers); and 4) number of cigarettes per day. Because former tobacco smoker status and age of tobacco smoking onset were measured only for individuals who had ever been tobacco smokers, they are not defined for the entire UKB sample, and we thus did not select these phenotypes.

For our remaining possibilities, we coded ever-tobacco smoker status as 1 if a respondent reported that they were a current or previous smoker and 0 if they reported never smoking or only smoking once or twice. We coded cigarettes per day as 0 if ever-smoking status was also 0; otherwise, we used the maximum number of reported past or current cigarettes (or pipes/cigars) consumed per day, normalized separately for males and females. Our cigarettes per day phenotype had a slightly higher phenotypic correlation with general risk tolerance in the first release of the UKB (0.098) than ever-smoker status (0.092). However, we concluded that the consistency of the ever smoker phenotype with previous GWAS literature overrides this slightly higher phenotypic correlation, and we therefore use this ever smoker phenotype as our final smoking behavior GWAS phenotype.

^a The normalized variable is the inverse normal cumulative distribution function (CDF) of the observations' percentile ranks.

^b <https://www.drinkaware.co.uk/alcohol-facts/alcoholic-drinks-units/latest-uk-alcohol-unit-guidance/>.

For our GWAS of ever smoker, we meta-analyzed the summary statistics from the UKB GWAS with those from the Tobacco, Alcohol and Genetics (TAG) Consortium³⁵ (the TAG consortium refers to the ever smoker phenotype as “smoking initiation”). In total, this meta-analysis includes 518,633 people from the meta-analyzed UK Biobank (444,598) and TAG (74,035) summary statistics.

Number of sexual partners: The potential phenotypes for assessing risky sexual behavior in the UKB include teenage conception (which we coded ourselves from available phenotypes related to age and pregnancy), age of first sexual intercourse, and lifetime number of sexual partners. Because we only defined teenage conception for females, we did not pursue this phenotype. Lifetime number of sexual partners has a much higher phenotypic correlation with general risk tolerance in the first release of the UKB (0.207) than does age of first sexual intercourse. We therefore use lifetime number of sexual partners (hereafter referred to as simply “number of sexual partners”) as our final risky sexual behavior GWAS phenotype. For this phenotype, respondents were asked, “About how many sexual partners have you had in your lifetime?” If respondents reported more than 99 lifetime sexual partners, they were asked to confirm their responses. We assigned a value of 0 to participants who reported having never had sex, and we again normalized this measure separately for males and females. In total, this GWAS includes 370,711 individuals in the UK Biobank.

First PC of risky behaviors: We performed a principal component analysis (PCA) with our four selected risky behaviors above and obtained the first principal component (PC) (see **Supplementary Table 1.4**). The first PC is the linear combination of the four risky behaviors that has the largest possible variance (among all possible linear combinations where the squares of the weights on the four risky behaviors sum to one). It can be interpreted as a general factor of risky behavior. As we describe below, we performed a GWAS of this first PC of risky behaviors, and we also examined the genetic correlation between this PC and general risk tolerance (see **Supplementary Table 7.1**). In total, this GWAS includes 315,894 people in the UK Biobank.

2 GWAS, quality control, and meta-analysis

2.1 Overview of main GWAS

All analyses were performed at the cohort level according to a pre-specified and publicly archived analysis plan³⁶. The original analysis plan was archived on February 4, 2016. For self-reported general risk tolerance, the analysis plan specified that the discovery GWAS would be conducted in the UKB and that the replication would be carried out in a meta-analysis of all other cohorts.

We updated the analysis plan on November 9, 2016 to include the analysis of the four risky behaviors and their first PC. For these phenotypes, the analysis plan specified that the GWAS would be conducted in the UKB. We did not attempt replication for these phenotypes. We updated the analysis plan a second time on July 10, 2017 to add the 23andMe cohort to the discovery GWAS of self-reported general risk tolerance. Two minor updates were made on August 7 and September 14, 2017 to specify that follow-up analyses (such as polygenic prediction) would be performed, whenever possible, on a meta-analysis combining the discovery and replication GWAS of general risk tolerance, and that we would add a GWAS of adventurousness alongside the primary GWAS of general risk tolerance and the other supplementary GWAS.

Cohorts other than the UKB could join this study by first supplying descriptive statistics and thereafter GWAS summary statistics from GWAS of self-reported general risk tolerance in November 2015 and December 2015, respectively. Two additional cohorts, Army STARRS and VHSS, joined the study later and provided descriptive and GWAS summary statistics in late 2016. Summary statistics were uploaded to a central, secure server and subsequently meta-analyzed. The lead PI of each cohort affirmed that the results contributed to the study were based on analyses approved by the local Research Ethics Committee and/or Institutional Review Board responsible for overseeing research. All participants provided written informed consent. We also obtained the descriptive and GWAS summary statistics from GWAS of self-reported general risk tolerance and adventurousness from 23andMe in late July 2017. An overview of the participating cohorts is reported in **Supplementary Table 1.1**.

The analysis plan instructed all cohorts to limit the analysis to individuals of European ancestry, to exclude individuals with missing covariates, to remove samples that displayed a SNP call rate of less than 95%, and to apply cohort-specific standard quality control filters before imputation. The cohort-specific standard quality control filters are reported in **Supplementary Table 2.1**. GWAS were limited to the 22 autosomes. The cohorts were required to provide unfiltered GWAS summary statistics including the following information for each SNP: chromosome and base-pair position, rsID, effect-coded allele, other allele, sample size per SNP, coefficient estimate (beta), standard error of the coefficient estimate, *P* value of the association uncorrected for genomic control, effect-coded allele frequency (EAF), imputation status, imputation quality, and Hardy-Weinberg equilibrium exact test *P* value for directly genotyped markers.

The analysis plan included power calculations assuming that 100,000 individuals in the UKB answered “Yes” (“cases”) to the general-risk-tolerance question, and 270,000 individuals answered “No” (“controls”). Under this assumption, our study would have 73% power to detect single nucleotide polymorphisms (SNPs) with a minor allele frequency (MAF) of 0.3 and an odds ratio of 1.05 with a genome-wide significance threshold of $P = 5 \times 10^{-8}$.

For general risk tolerance, the final sample size for the “discovery GWAS” meta-analysis of the UKB and 23andMe cohorts was 939,908 individuals. Replication was performed in a meta-analysis of 10 independent cohorts from seven studies totaling 35,445 individuals. We will henceforth refer to this meta-analysis of the 10 replication cohorts as the “replication GWAS.” The follow-up analyses we describe in the following **Supplementary Information sections** were performed with GWAS summary statistics from a meta-analysis combining the discovery and replication GWAS ($n = 975,353$), except where otherwise noted.

For adventurousness, GWAS summary statistics from 23andMe were analyzed ($n = 557,923$).

For three of the four risky behaviors, namely automobile speeding propensity ($n = 404,291$), drinks per week ($n = 414,343$), and number of sexual partners ($n = 370,711$), and for the first PC of the four risky behaviors ($n = 315,894$), GWAS were conducted in the UKB only, as specified in the updated analysis plan. For the remaining risky behavior, ever smoker, we meta-analyzed the summary statistics from the UKB GWAS ($n = 444,598$) with those from the Tobacco, Alcohol and Genetics (TAG) Consortium³⁵ ($n = 74,035$), leading to a total sample size of 518,633 (the TAG consortium refers to the ever smoker phenotype as "smoking initiation").

2.2 Genotyping and imputation

Genotyping^b was performed using a range of common, commercially available genotyping arrays. An overview of the genotyping and imputation procedure is provided in **Supplementary Table 2.1**. The participating cohorts were encouraged to use their standard quality-control protocols before imputation, as long as the applied filters satisfied the minimum requirements specified in the analysis plan (SNP call rate > 95%, HWE exact test P value > 10^{-6} , MAF > 1%). For the UKB, different filters were used, following ref.³⁷.

The cohorts, except for 23andMe, Army STARRS, BASE-II, UKB, and VHSS imputed markers using the 1000 Genomes phase 1 reference panel (March 2012 release version 3). 23andMe used the 1000 Genomes phase 1 (September 2013 haplotype release). Army STARRS used the 1000 Genomes phase 1 (August 2012 haplotype release). BASE-II used the more recent reference panel 1000 Genomes phase 3 (October 2014 haplotype release version 5). UKB used a customized reference panel based on the Haplotype Reference Consortium release 1.1 combined with the UK10K haplotype reference panel³⁸. VHSS used the Haplotype Reference Consortium release 1.1³⁹.

All genetic positions reported in this study are denoted with those of the Genome Reference Consortium’s human assembly 37 (GRCh37, sometimes referred to as the National Center for Biotechnology Information hg19).

2.3 Association analyses

Cohorts were encouraged to exclude individuals with SNP call rates less than 95%, with excessive autosomal heterozygosity, and with sex mismatch. Family-based cohorts were informed to control for family structure either with mixed linear modeling or with a procedure selecting only one individual in each pair that displayed relatedness greater than 5% in a genetic relatedness matrix.

^b The UKB genotype data was handled with QCtool, available at <http://www.well.ox.ac.uk/~gav/qctool/#overview>

The genome-wide association analysis performed in each cohort estimated the following regression for each SNP, as in Okbay *et al.* 2016¹⁶:

$$(1) \quad Y_i = \beta_0 + \beta_1 SNP_i + \mathbf{PC}_i \boldsymbol{\gamma} + \mathbf{X}_i \boldsymbol{\alpha} + \mathbf{C}_i \boldsymbol{\theta} + \epsilon_i,$$

where Y_i is the phenotype for individual i , SNP_i is the number of effect-coded alleles of the SNP^c, \mathbf{PC}_i is a vector of principal components of the genetic relatedness matrix after application of the pre-imputation filters described above, and \mathbf{X}_i is a vector of control variables. In all cohorts, \mathbf{X} included controls for sex and birth year. In most cohorts (including the 23andMe cohort), these included sex, birth year, birth year squared, birth year cubed, and the interactions between sex and the three birth-year variables; in the UKB, these included sex-specific birth year fixed effects. \mathbf{C}_i is a vector containing cohort-specific controls and technical covariates (such as dummy variables for genotyping array and genotyping batches) that are recommended in the analysis plan. All associations were performed with males and females pooled. A summary of the GWAS association models and control variables for each cohort is reported in **Supplementary Table 2.2**.

In practice, the phenotype was often residualized by first regressing the phenotype on the control variables, and the residualized phenotype was then regressed on the genotypes. This approach leads to almost identical results as estimating the full regression model directly while drastically reducing the computation time needed for the GWAS.

2.3.1 Linear mixed models in the UKB

The association analyses in the UKB were performed with linear mixed models (LMM) with the BOLT-LMM v2.2 software⁴⁰. The benefit of LMM is that the method accounts for cryptic relatedness and population structure, which allows the inclusion of related individuals in the sample, thereby yielding a larger sample size and greater statistical power. Using LMM is computationally intensive, and the BOLT algorithm is a new method that makes LMM analysis of hundreds of thousands of individuals computationally feasible. The method requires a set of SNPs to be included in the genetic variance component, and we included 483,680 directly genotyped bi-allelic autosomal SNPs with MAF > 0.005 and HWE P value > 10^{-6} . We included individuals based on self-reported ancestry, specifically those who self-reported to be of “white” ancestry (i.e., self-reported white, British, Irish, or any other white background). In addition, we limited the GWAS to individuals for whom the value of the first principal component of the genetic relatedness matrix was less than “0,” which identifies the cluster of individuals of European ancestry. We dropped individuals whose reported sex did not match their genetic sex, individuals with putative sex chromosome aneuploidy, individuals that did not pass the UKB internal genotype quality control, and individuals with missing values.

2.4 Main reference panel

The full release of the UK Biobank genetic data was imputed with haplotypes from the Haplotype Reference Consortium v1.1 (HRC) and the UK10K haplotype reference panel³⁸. A recommendation was communicated soon after the release of the genotype data in July 2017. It was recommended that only SNPs available in the HRC be used for analysis, because a subset of

^c For imputed SNPs we used best-guess data for samples that were imputed with IMPUTE²⁶⁷, and dosage data for samples that were imputed with MaCH/Minimac²⁶⁸.

variants imputed from the UK10K reference panel have wrongfully imputed genomic positions, while none of the HRC SNPs are affected. We therefore used the HRC v.1.1 as the reference panel for quality control of the GWAS summary statistics, and to determine the independence of significant SNPs. The following section describes our quality control of the HRC whole-genome sequence data (WGS) when constructing the reference panel. We will hereafter refer to the resulting reference panel as the “main reference panel.”

2.4.1 *Quality-control of the main reference panel*

The HRC haplotypes were downloaded from the European Genome-phenome Archive (EGA) on August 1, 2017. Strict internal quality control had already been applied to the WGS data, as described in-depth elsewhere³⁹, and we restricted the main reference panel to variants that passed all pre-applied genotype call filters (i.e., variants whose VCF FILTER status is “PASS”; these pre-applied filters included, among other filters, a filter to remove variants with minor allele count (MAC) ≤ 5 .) Before our internal QC the WGS data contained 40,405,506 autosomal and X-chromosome SNPs. The following protocol was restricted to the 39,131,579 autosomal SNPs because the pre-registered analysis plan restricts our analyses to the autosomes. The HRC reference panel does not include any structural variants, such as INDELs³⁹.

We performed a series of best-practice alignments of the WGS data for consistent and unique identification of variants⁴¹ with the open-source software BCFtools^d created by the Wellcome Trust Sanger Institute. Because PLINK cannot properly handle truly multi-allelic variants, we split multi-allelic variants into multiple bi-allelic variants. We then confirmed that all reference alleles and genomic positions matched the Genome Reference Consortium Human genome build 37 (GRCh37)⁴². To avoid issues with chromosomal positions mapping to multiple NCBI marker IDs (rsIDs), all rsIDs were removed, and all variants were given a unique identifier (henceforth referred to as the “unique ID”) in the form of chromosome, base-pair position, reference allele, and alternative allele, separated by colons (e.g., 1:123456:C:T). Using this format for variant IDs, together with the alignment to the reference genome GRCh37, ensures a unique representation of all SNPs and a lack of duplicate variants with switched reference alleles (e.g., 1:123456:C:T and 1:123456:T:C).

To avoid possible issues with inconsistencies with the UK10K haplotype reference in future work that may use that reference, we investigated possible strand and allele issues across the reference panels. By comparing the reference alleles and allele frequencies we found 24,394 variants with inconsistent alleles, and we decided to drop these from the main reference panel so that they would be removed during QC of the GWAS summary statistics.

We converted the VCF data to PLINK binary format with PLINK v.1.9b3.46⁴³, and we removed all multi-allelic variants (without retaining any of the multi-allelic variants coded as bi-allelic SNPs). Monomorphic SNPs (i.e., SNPs with MAF = 0) were kept in the reference panel as recommended³⁹. We thereafter excluded one member of each pair of individuals with genomic relatedness greater than 0.025 from the sample, which removed 4,917 individuals of the 22,691 individuals for whom data were available for all autosomes in the VCF data.

In summary, the main reference panel consists of 17,774 individuals and includes 38,889,224 bi-allelic autosomal SNPs that passed QC.

^d BCFtools can be downloaded here: <http://samtools.github.io/bcftools/bcftools.html>.

2.4.2 *rsID mapping*

Since rsIDs were removed from the SNPs to ensure unique identification, we created a map file so that each SNP could be assigned an rsID after meta-analysis, which is performed on the unique ID format described above (e.g., 1:123456:C:T). The map file contains the rsIDs from the HRC v1.1 sites list, and because we removed all multi-allelic variants there are no duplicate rsIDs.

2.5 UK Biobank genotyping arrays

2.5.1 *Combining data from the UK BiLEVE and the UK Biobank Axiom arrays*

The participants of the UKB were genotyped with two different but similar genotyping arrays^{38,44–46}. UKB participants who were enrolled in the UK BiLEVE study (a study of smoking behavior, lung function, and chronic obstructive pulmonary disease⁴⁴) were genotyped with the UK BiLEVE array ($n \sim 50,000$), and the remaining participants were genotyped with the UK Biobank Axiom array ($n \sim 400,000$). Henceforth, we will refer to these two sets of UK Biobank participants as the UK BiLEVE and the UKB Axiom cohorts.

While the UK Biobank (UKB) is a population-based study⁴⁵ collected during 2006–2010, the sample selection for the UK BiLEVE study began at a later stage, in 2012. The UK BiLEVE participants were selected from among the European-ancestry individuals in the UK Biobank⁴⁴, based on being in the “middles and extremes of the forced expiratory volume in 1 s (FEV₁) distribution among heavy smokers (mean 35 pack-years) and never smokers.”⁴⁴ Because the UKB Axiom cohort is the complement of the UK BiLEVE cohort, the UK Axiom cohort is also non-random, being under-sampled on heavy smokers and never smokers. It is thus only the complete UKB that is a population-based study without any particular sampling scheme based on lung function and smoking, while the UK BiLEVE and UKB Axiom cohorts are selected subsamples of the UKB.

We decided to analyze the UKB as a single cohort, rather than treating the UK BiLEVE and the UKB Axiom cohorts as two separate cohorts to be analyzed separately and then included in the meta-analysis as separate cohorts. (As indicated in **Supplementary Information section 2.3**, we included fixed effects to control for the genotyping arrays in the GWAS analyses.) Several factors led us to analyze the UKB as a single cohort. First, this allowed us to control for cryptic relatedness across the BiLEVE and Axiom samples with linear mixed models (LMM) with the BOLT-LMM v2.2 software⁴⁰. Analyzing the two cohorts separately would have necessitated dropping individuals who have relatives in the other cohort. Second, to our knowledge no published studies have analyzed the two cohorts separately. The pre-print of the UKB flagship paper³⁸, as well as many other recent large-scale GWAS^{47–49}, also analyzed the UKB as a single cohort.

During the revision stage, a Referee raised the point that our GWAS results could be sensitive to our decision of analyzing the UKB as a single cohort. Though we believe it is preferable to treat the UKB as a single cohort, it would be worrying if our results were sensitive to that decision. To verify that, we repeated our discovery GWAS of general risk tolerance and our GWAS of ever smoker, this time treating the UK BiLEVE and the UKB Axiom cohorts as two separate cohorts (and meta-analyzing the results). As we report in **Supplementary Information section 3.5**, the results barely changed.

2.5.1 Quality control of allele-frequency differences between the UK Biobank genotyping arrays

It was communicated that the first release of UKB data contained a small set of 65 autosomal SNPs that appeared to have flipped reference alleles contingent on the array. A subset of these had unfortunately been used during the imputation procedure. We already control for genotype array and batch during GWAS analyses, but as an additional QC step beyond excluding the 65 previously reported flipped SNPs, we investigated the allele frequencies across the arrays to be sure that our results were unaffected by artifacts from the genotyping procedure. It should be noted that the participants genotyped with the UK BiLEVE array were chosen based on lung function and smoking behavior⁴⁴, but the sample is in all other respects comparable to the rest of the UK Biobank⁵⁰.

We restricted the imputed genotype data to unrelated individuals of British ancestry to ensure that allele-frequency differences across the genotyping arrays would not be caused by differences in the proportion of ancestries or be affected by dependent observations. With PLINK⁴³, we calculated the allele frequencies contingent on the genotyping array for both the directly genotyped and imputed SNPs. Because our quality-control protocol, described in the next section, restricts the GWAS to SNPs with $MAF \geq 0.001$, we chose not to investigate SNPs with $MAF < 0.001$ (in the imputed genotype data) for allele-frequency differences between the UKB genotyping arrays. SNPs available on only one of the genotyping arrays and SNPs that were not available in our main reference panel were not considered in this investigation of allele-frequency differences.

For each SNP included in the investigation, three quantities measuring differences in allele frequencies were examined: the absolute value of the difference between the two arrays and the absolute value of the difference between the main reference panel and each of the arrays. We flagged a SNP as problematic if it fulfilled the following two conditions: (1) if the absolute value of the difference between the two arrays was greater than 0.25; and (2) if the absolute value of the difference between the main reference panel and at least one of the genotyping arrays was greater than 0.25. The comparison resulted in 600 flagged autosomal SNPs (including the 65 SNPs that were already reported as problematic) that were removed from the UKB summary statistics during QC in **Supplementary Information section 2.6.2**.

2.6 Description of major steps in quality-control (QC) analyses

For each cohort, we applied a stringent quality-control protocol based on the EasyQC software (version 9.2) developed by the GIANT consortium⁵¹, as well as additional steps developed by the Social Science Genetic Association Consortium^{16,18}. All issues raised during implementation of the protocol described below were resolved through iterations between the meta-analyst and the cohort analysts before any GWAS summary statistics were forwarded for meta-analysis.

2.6.1 Pre-QC verification and harmonization of GWAS summary statistics

All cohorts were asked to supply descriptive statistics and phenotype definitions according to the pre-specified analysis plan³⁶. The completeness of these documents was assessed as the first step of the quality control, together with examination of the uploaded GWAS summary statistics. All GWAS summary statistics were harmonized to ensure that the SNP identifier was in an admissible format (i.e. either an rsID, or in a format containing the chromosome, base pair (bp), and the two

alleles of the SNP), that the missing string operator was set to “NA,” and that all files had the same column delimiter.

2.6.2 *Filters applied before EasyQC protocol*

Following recommendations provided by the UK Biobank, we removed the 65 autosomal SNPs from the UKB that had been flagged as having incorrect annotation, together with the additional 535 SNPs that we flagged in section 1.5, before applying the EasyQC protocol described below.

Also, for cohorts imputed with the September or December 2013 haplotype release of the 1000 Genomes imputation reference panel, we removed 737 SNPs with incorrect strand alignment^e.

2.6.3 *EasyQC protocol*

The filters applied in the EasyQC software are explained below in chronological order of implementation. Note that the order of the filters does not influence the outcome of the cleaned GWAS summary statistics (although it affects at which specific filter a SNP is removed). The number of SNPs filtered at each step of the EasyQC protocol is reported in Panel A of **Supplementary Table 2.3**.

Step 1 in the EasyQC protocol filtered out SNPs for which either the effect-coded allele or the other allele has values different from “A,” “C,” “G,” or “T.” This step removed all structural variants such as INDELs.

Step 2 filtered out SNPs with missing values for one or more of the following variables: *P* value, an estimated effect size (beta) or its standard error, frequency of the reference allele, imputation status, and imputation accuracy (conditional on the SNP being imputed). This filter also removed SNPs with nonsense values outside of permissible ranges such as negative or infinite standard errors, nonsensical *P* values, allele frequencies greater than 1 or below 0, as well as imputation status not equal to 1 or 0.

The thresholds chosen for the filters applied in steps 3 to 5 are summarized in **Supplementary Table 2.4**. Step 3 filtered out SNPs with a MAF below 0.1% for the UKB and 23andMe cohorts and below 1% for all other cohorts; this effectively removed any SNPs that were monomorphic in the summary statistics. Step 4 excluded SNPs based on imputation accuracy with a threshold contingent on the cohort-specific imputation software (0.6 for MACH, 0.7 for IMPUTE, and 0.8 for PLINK). Step 5 filtered out directly genotyped SNPs with a Hardy-Weinberg equilibrium exact test *P* value below a threshold contingent on the cohort sample size. The applied thresholds were 10^{-3} if $n < 1,000$, 10^{-4} if $1,000 \leq n < 2,000$, and 10^{-5} if $2,000 \leq n < 10,000$.

Two additional filters were applied to ensure that only high-quality SNPs were being forwarded to the meta-analysis; step 6 removed SNP *j* if

^e The announcements are available on the webpage of IMPUTE2
https://mathgen.stats.ox.ac.uk/impute/data_download_1000G_1_integrated_SHAPEIT2_16-06-14.html and
https://mathgen.stats.ox.ac.uk/impute/data_download_1000G_phase1_integrated_SHAPEIT2_9-12-13.html.

$$(2) \quad \widehat{SE}_j > 1.4 \frac{\hat{\sigma}_Y}{\sqrt{2 \cdot n_j \cdot MAF_j \cdot (1 - MAF_j)}},$$

where $\hat{\sigma}_Y$ is the standard deviation of the phenotype, n_j is the sample size, SE_j is the standard error of the coefficient estimate for SNP j , and MAF_j is the minor allele frequency of SNP j . This filter eliminates SNPs whose coefficient estimates have standard errors that are more than ~40% larger than what would be expected given the sample size, the MAF of the SNP, and the standard deviation of the phenotype. The second additional filter, step 7, removes SNPs with coefficient estimates larger than what would correspond to an R^2 greater than 5%. We adapted the filter from Okbay *et al.*¹⁶ using an approximation to the R^2 : SNP j is dropped if

$$(3) \quad |\hat{\beta}_j| > \frac{\sqrt{0.05} \cdot \hat{\sigma}_Y}{\sqrt{2 \cdot MAF_j \cdot (1 - MAF_j)}}.$$

Step 8 filtered out duplicate SNPs (SNPs with identical NCBI build 37 (UCSC hg19) chromosome and base-pair positions). This was implemented after the chromosome and base-pair positions of the SNPs had been harmonized with the main reference panel described above.

Step 9 aligned the SNPs to the main reference panel to ensure that the effect-coded allele was the same for all SNPs across the cohorts. This step removed SNPs that were not available in the main reference panel as well as SNPs that displayed an allele mismatch when compared to the reference (e.g., a SNP with the alleles A and T in the GWAS summary statistics would be removed if the alleles according to the reference panel were A and G).

Step 10 removed SNPs that deviated from the main reference panel in terms of allele frequency. A SNP was removed if the absolute difference between its allele frequencies in the cohort's data and in the main reference panel was greater than 0.2. Step 10 was applied to all cohorts including the UKB (for the UKB, this filter was thus applied in addition to the filter described in **Supplementary Information section 2.5** and **Supplementary Information section 2.6.2**, the purpose of which was to avoid potential strand issues caused by the two different UKB genotyping arrays).

The output from the quality control was examined to see if any filters removed an unusual or unexpected number of SNPs. Some cohorts required iterations with the analysts to ensure that all possible errors were resolved. The number of SNPs filtered at each step of the final quality control iteration is reported in Panel A of **Supplementary Table 2.3** together with the estimated genomic inflation factor (λ_{GC}).

2.6.4 Visual inspection of diagnostic plots

Once low-quality SNPs were filtered out, the remaining SNPs were used to produce several diagnostic plots for each cohort, most of which are the standard output of the EasyQC software. Visual inspection of these plots enabled the identification of possible issues or errors in the GWAS summary statistics of each cohort; for a more thorough discussion we refer the interested reader to Winkler *et al.*⁵¹. For any potential issues observed in these plots, we contacted the cohort-specific

analyst and ensured that the observed issues were completely resolved. The following plots were examined:

Allele Frequency Plots (AF Plots): The AF plot contrasts the *observed* allele frequencies with the *expected* allele frequencies calculated according to the main reference panel, and the plot was created before step 10 of the EasyQC protocol. If the sample closely resembles the reference panel in terms of allele frequencies, then the SNPs should align in a diagonal with positive slope. This plot enables the analyst to detect deviations in ancestry from the reference as well as issues related to the alignment of the effect-coded allele. If the wrong effect-coded allele has been specified, then the AF plot shows a diagonal with negative slope.

P-Z Plots: Inspection of this plot shows if the reported P values are consistent with the reported coefficient estimates and their standard errors. One common problem observable with the P-Z plot is an erroneous column header in the GWAS summary statistics, such that the wrong column is used for either the beta estimates, standard errors or P values in the analysis.

Q-Q Plots: Inspection of Q-Q plots enables visualization of unaccounted-for stratification in the cohorts. No cohort displayed premature lift-off in the Q-Q plot associated with unaccounted-for stratification. The genomic inflation factors λ_{GC} are displayed in Panel A of **Supplementary Table 2.3**.

SE Plots: We plotted the observed standard errors (SE_j) of the coefficient estimates versus the standard errors expected given the MAF_j and the sample size n_j of a given SNP j , and the standard deviation of the phenotype $\hat{\sigma}_Y$ (which is equal to 1 if the phenotype has been standardized). This enables visual inspection to identify groups of outlier SNPs with regard to the observed standard error. The expected standard error was calculated according to the following formula:

$$(4) \quad \widehat{SE}_j \approx \frac{\hat{\sigma}_Y}{\sqrt{2 \cdot n_j \cdot MAF_j \cdot (1 - MAF_j)}}$$

All cohorts had to pass visual inspection as well as inspection of the number of excluded SNPs at each of the exclusion filters described in the previous subsection before being passed on for the meta-analysis.

2.7 Meta-analysis, adjustment of the standard errors, and test of heterogeneity of effect sizes across cohorts

2.7.1 Meta-analysis

Sample-size weighted meta-analysis of the cleaned cohort-level GWAS summary statistics were carried out using the METAL software⁵². We conducted four main meta-analyses: (1) we meta-analyzed the discovery GWAS combining the UKB and 23andMe cohorts; (2) we meta-analyzed the results of the 10 replication cohorts without the UKB and 23andMe discovery cohorts, to obtain our replication GWAS; (3) we meta-analyzed the results of the 10 replication cohorts together with those of the UKB and 23andMe discovery cohorts for the follow-up analyses that use GWAS summary statistics; and (4) we meta-analyzed the results from our UKB GWAS of ever smoker with those from the TAG Consortium³⁵. No meta-analyses were conducted for the five other

supplementary GWAS, because data for these GWAS each came from only one cohort (either the UKB or the 23andMe cohort).

All meta-analyses were performed with the unique ID format as the identifier of each SNP (e.g., 1:123456:C:T)¹⁸. All meta-analyses were restricted to SNPs with a sample size greater than half of the maximum sample size across all the SNPs in the GWAS. Thus, because the discovery GWAS of general risk tolerance consists only of the 23andMe and UKB cohorts and because the 23andMe cohort is slightly larger than the UKB, all 9,284,738 SNPs available in the 23andMe cohort (and no other SNPs) were analyzed in the discovery GWAS of general risk tolerance. Of these 9,284,738 SNPs, 8,989,321 are available in both the UKB and the 23andMe cohorts and have a sample size of 931,651 or 939,908^f, and 295,417 are available only in the 23andMe cohort and have a sample size of 500,525 or 508,782^g. Of the 124 general-risk-tolerance lead SNPs we report below in **Supplementary Information section 3.3**, all but one (rs13251864) are present in both the 23andMe and UKB cohorts. For the replication GWAS of general risk tolerance, 6,986,015 SNPs were analyzed; 9,339,358 SNPs were analyzed in the GWAS of adventurousness; and ~11,515,000 SNPs were analyzed in the GWAS of the four risky behaviors and their first PC. Panel **B** of **Supplementary Table 2.3** reports the number of SNPs for all the main GWAS.

2.7.2 *Adjustment of the standard errors*

Instead of applying genomic control with the over-conservative λ_{GC} , we inflated the standard errors by the square root of the estimated intercept from an LD Score regression. This procedure allows us to correct only for inflation of test statistics caused by population stratification and other confounding factors rather than polygenicity⁵³.

For the discovery and replication GWAS of general risk tolerance and for the GWAS of ever smoker—all of which involved meta-analyses of cohort-level data—we only inflated the meta-level standard errors (i.e., we did not inflate the cohort-level standard errors before the meta-analysis). Likewise, for the meta-analysis of the discovery and replication GWAS for the follow-up analyses, we only inflated the meta-level standard errors. We also inflated the standard errors of the other supplementary GWAS.

In practice, for a given meta-analysis, the METAL software⁵² outputs the SNPs' meta-analyzed z -statistics, deflated by the square root of the estimated intercept from an LD Score regression. We use SNP j 's GWAS sample size n_j and minor allele frequency MAF_j , as well as the phenotype's standard deviation $\hat{\sigma}_y$, to approximate the inflated standard error of our estimate of SNP j 's effect size:

$$\widehat{SE}_j \approx \sqrt{Intercept} \cdot \frac{\hat{\sigma}_y}{\sqrt{2 \cdot n_j \cdot MAF_j (1 - MAF_j)}},$$

where $\sqrt{Intercept}$ is the square-root of the LD Score intercept used to deflate the z -statistic in the meta-analysis.

^f 8,949,622 SNPs have a sample size of 939,908 and 39,699 SNPs have a sample size of 931,651.

^g 290,259 SNPs have a sample size of 508,782 and 5,158 SNPs have a sample size of 500,525.

We then used SNP j 's deflated z -statistics \hat{z}_j to approximate SNP j 's effect size as $\hat{\beta}_j \approx \hat{z}_j \cdot \widehat{SE}_j$ ^h. Since the general-risk-tolerance phenotype is not measured in natural units, and since its standard deviation differs across cohorts, we normalize it to have a standard deviation of one when we estimate the SNPs' effect sizes and standard errors. For consistency, we make the same assumption when approximating the SNPs' effect sizes and standard errors for the risky behaviors and their first PC. Hence, our estimated effect sizes (the $\hat{\beta}_j$'s) are expressed in standard-deviation units of the phenotype per effect-coded allele; we hereafter refer to this as a “standardized beta,” although it is only standardized in terms of standard deviation units of the phenotype (and not with respect to the genotype). The standard deviations of the phenotypes, as originally measured, are reported in **Supplementary Table 1.2**.

The coefficient of determination of SNP j is approximated as⁵⁴:

$$(5) \quad R_j^2 \approx \frac{2 \cdot MAF_j(1 - MAF_j) \cdot \hat{\beta}_j^2}{\hat{\sigma}_y^2}.$$

2.7.3 *Evaluation of effect-size heterogeneity across cohorts for general risk tolerance*

Following a Referee's suggestion, we computed Cochran's Q statistic for the lead SNPs of our discovery GWAS of general risk tolerance, to evaluate the heterogeneity of our estimates across the 23andMe and UKB cohorts. (We note, however, that the power of Cochran's Q test is limited in our setting^{55,56}, because the discovery meta-analysis consists of only two cohort.) In addition to examining the P values of Cochran's Q test for each lead SNP (after Bonferroni correction for the number of lead SNPs), we generated an omnibus test statistic for heterogeneity by summing the Cochran Q statistics across all lead SNPs⁵⁷. Because there are two cohorts, the Q statistic for each lead SNP has a χ^2 distribution with one degree of freedom. The sum of these Q statistics is therefore (approximately) χ^2 -distributed with the number of degrees of freedom being equal to the number of lead SNPs. We report the results in **Supplementary Information section 3.3.2**.

2.8 **Approximately independent lead SNPs, loci, and conditional analysis**

2.8.1 *Approximately independent lead SNPs*

To identify approximately independent genome-wide significant “lead SNPs”, we used PLINK⁴³ to apply a “clumping algorithm” to the GWAS results. (We define a SNP as “genome-wide significant” if its GWAS P value is less than 5×10^{-8} .) Our clumping algorithm uses four parameters: a primary P value threshold (5×10^{-8}), a secondary P value threshold (1×10^{-4}), an r^2 threshold (0.1), and a SNP window defined in kilobases (1,000,000 kb). First, the SNP with the lowest P value (less than the primary P value threshold) is taken as the “lead SNP” in the first clump, and the first clump is formed by all SNPs with a P value smaller than the secondary P value thresholdⁱ, with an r^2 greater than 0.1 with the clump's lead SNP, and within a distance less than the SNP window from the lead SNP. (We used a very wide SNP window of 1,000,000 kb, which

^h Since \hat{z}_j is deflated and \widehat{SE}_j is inflated by the square root of the intercept from the LD score regression, $\hat{\beta}_j$ is neither deflated nor inflated.

ⁱ The secondary P value threshold lowers the computational effort by allowing the algorithm to ignore SNPs with large P values.

effectively makes the r^2 and P value thresholds the only binding parameters for the PLINK clumping algorithm.) Next, the SNP with the second lowest P value (less than the primary P value threshold) outside the first clump becomes the lead SNP of the second clump, and the second clump is created analogously but using only the SNPs outside of the first clump. This process continues until every SNP with a P value less than the primary P value threshold is either defined as the lead SNP of a clump or clumped with another lead SNP. The r^2 was calculated with the main reference panel. Thus, a “lead SNP” is the most significant genome-wide significant SNP in an approximately independent clump, and a lead SNP cannot be in the clump of another lead SNP for the same phenotype.

2.8.2 *Definition of non-overlapping, continuous genomic loci*

For the purpose of defining non-overlapping, continuous genomic loci, we followed Ripke *et al.*⁵⁸. Ripke *et al.* defined a locus as “the physical region containing all SNPs correlated at $r^2 > 0.6$ with [one of the lead] SNPs”, and merged associated loci within 250kb of each other into a single locus. For each of the seven GWAS, we followed this definition and created a set of loci. We report the identified loci in **Supplementary Information section 3**, and the loci are listed **Supplementary Table 3.1** and **3.2**. In that section, we also report the loci we obtained after pooling all the loci from across the seven GWAS and merging loci within 250kb of each other; those loci are listed in **Supplementary Table 3.3**.

2.8.3 *Conditional and joint multiple-SNP (COJO) analysis with GCTA*

Because we consider approximately independent (pairwise $r^2 < 0.1$) lead SNPs and loci (rather than fully independent lead SNPs and loci), some of our lead SNPs could in principle be secondary associations that are driven by their LD with extremely strong primary associations. We thus performed conditional and joint multiple-SNP (COJO) analysis⁵⁹ with GCTA. For each of the seven GWAS, we restricted the analysis to the set of SNPs that (1) pass all GWAS quality control filters, and (2) are located within the loci of the phenotype (which includes all the lead SNPs). We analyzed the summary statistics using the stepwise model-selection algorithm detailed in the original COJO publication⁵⁹. The analysis requires two input parameters: (1) the distance in kb at which perfect linkage equilibrium ($r^2 = 0$) is assumed, and (2) an r^2 threshold that prevents the stepwise model selection from adding SNPs that are highly correlated with a previously selected SNP. We used the default parameters, which assume perfect linkage equilibrium for SNPs separated by 10 Mb and which do not add SNPs in strong LD ($r^2 > 0.9$) with an already selected SNP. The COJO analysis was performed with LD estimated in our main reference panel (described in **Supplementary Information section 2.4**). We report the results of the COJO analysis in **Supplementary Information section 3**.

As we discuss in **Supplementary Information section 3.6**, we also conducted a multiple regression analysis with individual-level data from the UKB. In that analysis, for each phenotype (except adventurousness, for which there is no UKB data), for each chromosome we regressed the phenotype on all the phenotype’s lead SNPs located on the chromosome (and on control variables). The results were consistent with those of the COJO conditional analysis.

2.9 Check for long-range LD regions, candidate inversions, and 1000 Genomes structural variants

2.9.1 Check for long-range LD regions

We investigated if the lead SNPs were located in larger structural variation in the form of long-range LD regions, by using a set of long-range LD regions from Price *et al.*⁶⁰. Price *et al.* identified 24 long-range LD regions from 327 European-ancestry individuals, which replicated in two independent samples comprising 1593 European-ancestry Americans and 3004 British individuals. We lifted the genomic positions of the long-range LD regions to build 37 (GRCh37) with the UCSC genome-annotation lift-over tool^j, and there were nine long-range LD regions that could not be lifted due to non-overlapping genome sequences or ambiguous mapping across the builds. Hence, the combined map contains 15 long-range LD regions that have non-ambiguous genomic positions available in build 37. These range from ~2.5 to ~8 Mb in genomic size.

We checked if each lead SNP from the GWAS was within a long-range LD region, or within 250 bp from the breakpoints of such a region. The results are reported in **Supplementary Table 3.1** and **Supplementary Table 3.2**.

2.9.2 Check for candidate inversions

We also investigated if the lead SNPs were located in larger structural variation in the form of candidate inversions, by using a list of genomic segments highly prone to inversion polymorphisms from an unpublished resource by Gonzalez, J.R. & Esko, T., (2017, unpublished). The genomic segments highly prone to inversion polymorphisms were identified based on the knowledge that submicroscopic human inversions are typically flanked by highly homologous flanking repeats⁶¹, which predisposes their occurrence by non-allelic homologous recombination. Therefore, a set of segments was selected that may be prone to submicroscopic inversions, consisting of all single copy segments in the Genome Reference Consortium's human reference sequence build 36 (GRCh36) between 0.1 and 8 Mb in length, and flanked by segmental duplications with 90% identity (across the flanking duplications). In total, there were 173 segments that met these criteria and that were thus considered as genomic segments highly prone to inversions. As detectable traces of inversions in SNPs depend on many factors—such as being frequent, ancient and nonrecurring—we tested whether the segments showed inversion patterns in any of two different SNP datasets. First, 69 (40%) of the segments overlapped with the inversions that Caceres *et al.*⁶² obtained in the phased genotypes of CEU individuals from the HapMap III project. Second, inversion-like haplotypes⁶³ were inferred in a subsample of 882 Estonians for which gene expression data was available in peripheral blood. In this case 65 (38%) of the 173 segments were significantly associated with the expression of single copy genes within the segment. In total 104 (60%) of the 173 segments showed an inversion signal, indicating their predisposition for inversion occurrence.

We lifted the genomic positions of the genomic segments highly prone to inversion polymorphisms to build 37 (GRCh37) with the UCSC genome-annotation lift-over tool^k, and there were 19 genomic segments that could not be lifted due to non-overlapping genome sequences or ambiguous

^j The lift-over tool is available here: <https://genome.ucsc.edu/cgi-bin/hgLiftOver>

^k The lift-over tool is available here: <https://genome.ucsc.edu/cgi-bin/hgLiftOver>

mapping across the builds. Hence, the combined map contains 154 segments prone to inversion polymorphisms (hereafter referred to as “the 154 candidate inversions”), that have non-ambiguous genomic positions available in build 37. These range from ~500 kb to ~8 Mb in genomic size.

We checked if each lead SNP from the GWAS was within such a candidate inversion, or within 250 bp from the breakpoints of such a candidate inversion. The results are reported in **Supplementary Table 3.1** and **Supplementary Table 3.2**.

2.9.3 Check for 1000 Genomes structural variants

Sudmant *et al.* (2015)⁶⁴ called and classified a large number of structural variants (SVs) with the final version of the 1000 Genomes Project phase 3 reference panel. They have released an integrated map of 37,250 smaller structural variants, together with enhanced resolution of the size and breakpoint compared to previous publications, and we hereafter refer to these as the “1000G structural variants.” The structural variants range from 1 bp to ~445 kb in genomic size. The smallest variants are generally insertions of 1 bp (~6,900 variants), and the larger variants are generally deletions larger than 50 bp. The majority of these structural variants are in LD with proximate SNPs, and we therefore checked if any of the lead SNPs from our main GWAS, and SNPs in strong LD with those lead SNPs, were located within the start and end positions of any of the 37,250 structural variants. We defined strong LD as an r^2 greater than 0.8, which is the definition used by the 1000 Genomes Project Consortium⁶⁴. The SNPs in LD were extracted using PLINK⁴³ and the main reference panel. We report the results in **Supplementary Table 3.1** and **Supplementary Table 3.2**.

2.10 Investigation of the novelty of our GWAS associations

To investigate the novelty of our GWAS associations, we performed lookups of our lead SNPs (and the SNPs in LD with the lead SNPs, $r^2 > 0.1$) in the NHGRI-EBI GWAS Catalog database (revision 2017-08-15)⁶⁵ of genome-wide significant associations from previous GWAS. We also looked up our lead SNPs (and the SNPs in LD, $r^2 > 0.1$) in some recent GWAS articles that have not been catalogued in the NHGRI-EBI GWAS Catalog database. The NHGRI-EBI GWAS Catalog is a resource that aims to catalogue all associations reported in published GWAS.

For general risk tolerance, we performed a search with the term “risk” in the index of phenotypes, and we did not find any previous studies on general risk tolerance in the Catalog. We know of one previous study that identified one independent genome-wide significant association with general risk tolerance, and of one concurrent study that identified a second genome-wide significant association, both using the first UKB data release^{66,67}; the authors of ref.⁶⁶ referred to the phenotype as “risk-taking propensity,” and the authors of ref.⁶⁷ referred to it as “risk-taking behavior.” The first genome-wide significant association is replicated in an online publication published in advance⁶⁸. We added the first of these two studies (i.e., Day *et al.*⁶⁶) to our investigation of the novelty of our general-risk-tolerance lead SNPs, and the second we consider concurrent. We also note that, in **Supplementary Information section 11.1**, we report the results of a literature search of association studies of risk tolerance; that literature search identified no previously reported genome-wide significant associations.

The phenotypes drinks per week, ever smoker, and number of sexual partners (or related phenotypes such as alcoholism and age at first sex), were available in the NHGRI-EBI GWAS Catalog database (revision 2017-08-15). Since the GWAS Catalog is not always up-to-date, we

additionally performed a literature search for genome-wide significant findings that might not yet have been included in the Catalog. We searched the Pubmed literature database on March 6 2017 and September 13 2017, for the term “genome-wide association study” together with each of the terms “alcohol,” “sexual,” and “smoking” individually. We screened the abstracts and compared the resulting articles with the article list of the GWAS catalog. In addition to what was already reported in the GWAS Catalog (revision 2017-08-15), we found three additional studies with genome-wide significant findings on alcohol consumption, two additional studies on smoking, and no additional studies on sexual behaviors.

To our knowledge, this is the first GWAS of adventurousness, automobile speeding propensity, and of the first PC of the four risky behaviors; unsurprisingly, we could not find previous GWAS on any of these phenotypes in the NHGRI-EBI GWAS Catalog database (revision 2017-08-15).

2.11 GWAS catalog lookup

We investigated whether the lead SNPs from our discovery GWAS of general risk tolerance and from our supplementary GWAS have previously been associated at genome-wide significance with any phenotypes in the NHGRI-EBI GWAS Catalog database⁶⁵ (revision 2017-08-15). We query the GWAS Catalog with our list of lead SNPs, and the SNPs in LD with a lead SNP ($r^2 > 0.6$).

Since the NGRI-EBI GWAS Catalog is not always up-to-date with results from the most recent publications (especially in non-peer reviewed outlets such as BioRxiv), we also queried the summary statistics of the most recently published GWAS on attention deficit hyperactivity disorder⁶⁹, autism spectrum disorder⁷⁰, and anorexia nervosa¹. We also queried the genome-wide significant findings from the additional GWAS on alcohol intake and smoking, which we found in addition to the GWAS Catalog, as detailed in the previous section. We queried the additional GWAS on alcohol intake and smoking because they contained at least one genome-wide significant result; because their results were publicly available; and because the meta-analysis that produced them did not include the UK Biobank (that comprise our discovery sample together with 23andMe).

We perform these lookups because the existence of SNPs and genes associated with both one of our studied phenotypes and another phenotype can point to a common genetic etiology. However, it is important to note that two phenotypes that share a genetic locus do not necessarily have to share the same causal *variant* at that locus due to the widespread LD that characterizes the human genome. Moreover, even if two phenotypes do share a single causal variant, they do not have to share general underlying genetic etiologies. For instance, recent work²⁴ has shown that two types of autoimmune diseases (rheumatoid arthritis and ulcerative colitis/Crohn’s disease) that are known to share risk loci are not genetically correlated at a genome-wide level. Here, the reason was the lack of an overall directional trend: some risk alleles for one disease were also risk alleles for the other disease, but some alleles that were *protective* for the one disease were risk alleles for the other. This resulted in a near-zero correlation at the genome-wide level. Thus, we emphasize that the current lookup does not make it possible to determine *etiological* overlap, but only hints

¹ The Psychiatric Genomics Consortium’s GWAS summary statistics for attention deficit hyperactivity disorder, autism spectrum disorder, and anorexia nervosa (referred to as “ED,” i.e. eating disorder) are publicly available and can be downloaded here: <https://www.med.unc.edu/pgc/results-and-downloads>.

at overlapping loci, between general risk tolerance or the supplementary GWAS phenotypes, and the phenotypes reported in the GWAS Catalog.

2.12 Cross-lookup of GWAS results

We performed cross-lookups of the lead SNPs across our discovery GWAS of our primary and supplementary phenotypes. Specifically, for each lead SNP in each of the GWAS, we checked if the SNP is in LD (with an r^2 greater than 0.1) with lead SNPs in the other GWAS. LD was calculated using PLINK⁴³ and the main reference panel. The results of the cross-lookups are reported in **Supplementary Table 3.1** and **Supplementary Table 3.2**. As we describe above, we also investigated if there were any long-range LD regions or candidate inversions that contained lead SNPs for multiple GWAS.

2.13 Gene annotation

We annotated the lead SNPs with gene information using the National Institute of Health (NIH) National Center for Biotechnology Information (NCBI) gene ontology database (version 2016-05-25)^m. As the general rule, a SNP was annotated to its most proximate gene. If a SNP was located between two genes, then we compared the distance to the end coordinate of the gene upstream with the distance to the start coordinate of the gene downstream to find the most proximate gene. If a SNP was located within multiple overlapping genes, then the SNP was annotated to the gene with the most proximate start coordinate. This means that all lead SNPs were annotated to a single gene. The approach roughly partitions the SNPs throughout the genome into separate genomic segments. The annotations are displayed in **Supplementary Table 3.1** and **Supplementary Table 3.2**, where we also indicate if a lead SNP is located within or outside the gene's start and end coordinates. We also checked if there were genes to which lead SNPs from multiple GWAS were annotated (the results are reported in **Supplementary Information section 3.2**).

^m The NCBI gene ontology database is available here:
ftp://ftp.ncbi.nlm.nih.gov/gene/DATA/GENE_INFO/Mammalia/.

3 GWAS results

In this section, we report, compare, and discuss the results of our seven main GWAS (of our primary phenotype—self-reported general risk tolerance—and of our six supplementary phenotypes—adventurousness, the four risky behaviors, and their first PC). In **Supplementary Information sections 3.1 and 3.2**, we present a summary of our results, and we also describe several notable genomic regions that contain lead SNPs for all or most of our main GWAS. **Supplementary Information sections 3.3 and 3.4** contain a more detailed description of the results of our discovery GWAS of general risk tolerance and the results of our six other main GWAS, respectively. Throughout, we place particular emphasis on long-range LD regions and candidate inversions (defined in **Supplementary Information section 2.9**) that contain lead SNPs for all or most of our GWAS. (We focus on these because, as we explain below, very few genomic blocks outside of these regions or candidate inversions contain lead SNPs shared across most of our GWAS.)

3.1 Summary of the GWAS results

We identified a total of 864 “lead associations” across our seven main GWAS, where we define a lead association as a lead SNP in one of the GWAS: 124 lead SNPs associated with general risk tolerance, 167 with adventurousness, 42 with automobile speeding propensity, 85 with drinks per week, 223 with ever smoker, 117 with number of sexual partnersⁿ, and 106 with the first PC of the four risky behaviors. **Supplementary Information section 5** reports the results of our successful attempt to replicate the associations of our 124 lead SNPs with general risk tolerance. We did not attempt replication of the results of our six supplementary GWAS in independent data, because we did not have access to such data for the six supplementary phenotypes. However, as we report in **Supplementary Information section 5**, we calculated the “maxFDR”⁷¹, an upper bound on the false discovery rate (FDR), for each GWAS. The maxFDR estimates were low across all GWAS (the highest estimate was 1.22×10^{-3} , for automobile speeding propensity), thus providing reassurance about the robustness of the lead associations. **Supplementary Tables 3.1-3.2** report the detailed association results, and regional association plots are provided with the online supplementary materials

To the best of our knowledge, 852 of the 864 lead associations are novel. We were able to replicate the only previously published⁶⁶ genome-wide significant association with general risk tolerance, located within *CADM2* on chromosome 3, that was also replicated in ref.⁶⁸ and in a concurrent study⁶⁷. We replicated another genome-wide significant association with general risk tolerance from the concurrent study⁶⁷, located in proximity to the HLA-complex on chromosome 6. We also replicated the TAG Consortium’s previous association with ever smoker in the gene *NCAMI*³⁵.

The detailed results of the cross-lookup of our GWAS results (the investigation of whether the lead SNPs of each of our GWAS are in LD, defined as a r^2 greater than 0.1, with the lead SNPs of our other GWAS; **Supplementary Information section 2.12**) are reported in **Supplementary Tables 3.1 and 3.2**. In total, we identified 864 lead SNPs across the GWAS of our primary and supplementary phenotypes. Of these 864 lead SNPs, 34 exact lead SNPs are counted twice by

ⁿ Our baseline GWAS protocol (described in **Supplementary Information section 2**) identified 118 number-of-sexual-partners lead SNPs, but we excluded one of these from the count because we suspected it may not have been properly genotyped (see **Supplementary Information section 3.4.5** for details).

being shared by two phenotypes (there are no exact lead SNPs shared across more than two phenotypes; thus, in total there are 830 unique lead SNPs identified across the phenotypes). 441 of the 864 lead SNPs are in weak LD (pairwise $r^2 > 0.1$) with a lead SNP for at least one of the other GWAS.

Applying our locus definition, we identified 99 loci associated with general risk tolerance, 137 loci associated with adventurousness, 36 loci associated with automobile speeding propensity, 62 loci associated with drinks per week, 183 loci associated with ever smoker, 97 loci associated with number of sexual partners^o, and 89 loci associated with the first PC of the four risky behaviors. We will refer to these loci as the 703 “locus associations”, where a locus association is defined as a locus in one of our seven GWAS (see **Supplementary Information section 2.8** for our locus definition). LocusZoom plots of the 703 locus associations are provided with the online supplementary materials⁷². Pooling the loci corresponding to the 703 locus associations from across the seven GWAS and merging loci within 250kb of each other yielded 444 distinct loci; those loci are listed in **Supplementary Table 3.3**.

We supplemented those analyses with a conditional and joint multiple-SNP (COJO) analysis⁵⁹ with GCTA (**Supplementary Information section 2.8.3**). The COJO analysis identified a total of 655 conditional associations across all seven GWAS.

The 864 lead associations and 444 loci across the seven GWAS were obtained by using the standard genome-wide significance P value threshold of 5×10^{-8} to identify the lead SNPs for each phenotype. If we instead consider the seven GWAS jointly and use a Bonferroni-corrected P value threshold of 7.1×10^{-9} ($= 5 \times 10^{-8}/7$), across the seven GWAS we obtain 566 lead SNPs (instead of 864), and 464 locus associations (instead of 703). Pooling and merging the 464 locus associations yielded 304 distinct loci (instead of 444), and 505 conditional associations (instead of 655).

In sum, across our seven GWAS we identified 864 lead SNPs, 703 locus associations, and 655 conditional associations, and the loci corresponding to the 703 locus associations span 444 distinct loci.

The NHGRI-EBI GWAS Catalog database⁶⁵ lookup of general-risk-tolerance lead SNPs resulted in 61 overlapping associations distributed across 27 lead SNPs, and the lookup of our six other main GWAS resulted in 939 overlaps distributed across 130 lead SNPs. Notably, for all our main phenotypes we find overlaps with schizophrenia, cognitive performance, and information processing speed. We report the results of this lookup in **Supplementary Table 3.4**.

3.2 Summary of genetic overlap across our main GWAS

3.2.1 *Summary and discussion of notable genomic regions*

Five genomic regions stand out because they contain lead SNPs for all or most of our seven phenotypes. (Several of these regions contain multiple lead SNPs associated with one of the seven phenotypes. Most of the multiple lead SNPs within these regions are not conditional associations, but typically at least one and sometimes two of the lead SNPs within each region are conditional associations. Therefore, the exact numbers of lead SNPs within these regions should be interpreted

^o Our baseline GWAS protocol (described in **Supplementary Information section 2**) identified 98 number-of-sexual-partners loci, but we excluded one of these from the count because we suspected one of the lead SNP may not have been properly genotyped (see **Supplementary Information section 3.4.5** for details).

with caution.) Two of these regions are among the 15 long-range LD regions identified by Price *et al.*⁶⁰, and the other three are among the 154 genomic segments deemed highly prone to inversion polymorphisms (i.e., the 154 “candidate inversions”; both the 15 long-range LD regions and the 154 candidate inversions are described in **Supplementary Information section 2.9**). One long-range LD region (chromosome 3, ~83.4 to 86.9 Mb, **Extended Data Fig. 3.4**) and one candidate inversion (chromosome 18, ~49.1 to 55.5 Mb, **Extended Data Fig. 3.4**) each contain lead SNPs and conditional associations for all of our seven GWAS phenotypes; the other three regions each contain lead SNPs and conditional associations for general risk tolerance and for four or five of our six supplementary GWAS.

The first notable region is the long-range LD region on chromosome 3 (~83.4 to 86.9 Mb, **Extended Data Fig. 3.4**), which contains lead SNPs and conditional associations for all our GWAS. The only gene within the long-range LD region is *CADM2*, and there are few other genes in proximity (only *VGLL3* is within 250 kb of the breakpoints, ~69.8 kb downstream). The relatively large *CADM2* gene (~85.0 to 86.2 Mb) covers ~1.2 Mb of the ~3.5 Mb long-range LD region and contains our strongest association with general risk tolerance (rs993137, $P = 2.14 \times 10^{-40}$). The Bonferroni-corrected P -value of *CADM2* in the MAGMA gene analysis is $P = 1.09 \times 10^{-50}$ (**Supplementary Table 12.3**), consistent with the GWAS results. A recent study⁶⁸ reports that *CADM2* contains a replicated association with general risk tolerance, and that study also reports suggestive associations between SNPs in *CADM2* and different measures of personality. As we discuss in **Supplementary Information section 12**, *CADM2* “encodes a member of the synaptic cell adhesion molecule 1 (SynCAM) family which belongs to the immunoglobulin (Ig) superfamily”⁷³, and it is related to synapse formation⁷⁴ and brain plasticity⁷⁵. *CADM2* is overexpressed in the brain, and in particular in the frontal cortex, according to GTEx⁷⁶. The GWAS Catalog database⁶⁵ reports genome-wide significant associations in the long-range LD region with many phenotypes, including age at menarche, BMI, educational attainment, and information processing speed. Most of the GWAS Catalog database⁶⁵ associations within the long-range LD region (~83.4 to 86.9 Mb on chromosome 3) are annotated to the gene *CADM2*.

The second notable region is a candidate inversion located on chromosome 18 (~49.1 to 55.5 Mb, **Extended Data Fig. 3.4**). The candidate inversion contains lead SNPs and conditional associations for all phenotypes. Within the candidate inversion there are previous genome-wide associations in the GWAS Catalog database⁶⁵ with traits such as autism spectrum disorder, ADHD, depression, educational attainment, schizophrenia, and subcortical brain region volumes. The candidate inversion contains ~20 genes, and the MAGMA gene analysis resulted in one significant gene after Bonferroni correction—*TCF4* (Bonferroni-corrected $P = 5.51 \times 10^{-9}$, **Supplementary Table 12.3**). *TCF4* is interesting because it is known to play an important role in nervous system development⁷³. *De novo* mutations in *TCF4*⁷⁷ are known to cause the rare Pitt-Hopkins syndrome⁷⁸, with few described cases in the medical literature⁷⁹. The syndrome is characterized by distinct facial features, intellectual disability, delayed motor skills, and epilepsy, among many other symptoms^{77–79}. The GWAS Catalog database⁶⁵ reports genome-wide significant associations mapped to *TCF4* with schizophrenia, and *TCF4* has been hypothesized to be involved in more neuropsychiatric phenotypes⁸⁰. This observation is consistent with the non-zero genetic correlations that we estimated with bivariate LD Score regression between general risk tolerance and many neuropsychiatric disorders (**Fig. 2, Supplementary Information section 7**). As we describe below, all phenotypes have lead SNPs annotated to *TCF4* except ever smoker, for which there are three lead SNPs (of which two are conditional associations) within the candidate inversion annotated to the proximate genes *DCC* and *TXNLI*.

The third notable genomic region is the long-range LD region located on chromosome 6 (~25.3 to 33.4 Mb, **Extended Data Fig. 3.4**). The region covers the HLA-complex⁷³ (~29.6 to 33.1 Mb on chromosome 6⁸¹), and contains lead SNPs and conditional associations for all phenotypes except drinks per week (for which we identified a suggestive association, with a $P = 3.83 \times 10^{-7}$). There are at least 250 genes in the region, and the MAGMA gene analysis resulted in ~30 significant genes after Bonferroni-correction (however, none of the actual HLA genes were significant; **Supplementary Table 12.3**). The GWAS Catalog database⁶⁵ reports more than a thousand genome-wide significant associations in the region. The associations relate to hundreds of traits, including alcohol consumption, Alzheimer's disease, autism spectrum disorder, educational attainment, and schizophrenia. The HLA-complex contains thousands of SNP associations in the GWAS Catalog database⁶⁵, and those relate to hundreds of traits, including alcohol consumption, Alzheimer's disease, autism spectrum disorder, and educational attainment. The HLA genes encode the major histocompatibility complex (MHC) proteins, whose main function is to alert cells of the immune system about infection from pathogens⁸². The region is known to be highly polygenic, and its high rates of recombination lead to a large number of possible haplotypes in the population. For certain classes of MHC proteins there are more than 1,000 alleles in the human population, and most individuals are therefore heterozygous. However, there is also a small subset of proteins encoded for which the coding alleles are practically monomorphic, and this is probably caused by certain constraints on the viability of the variability for these specific protein chains. The HLA genes have been found to be under strong selection^p. The constant struggle to counter pathogens results in selection pressures that favor polymorphisms that are able to provide protection.

The fourth notable genomic region is the candidate inversion on chromosome 7 (~124.6 to 132.7 Mb; **Extended Data Fig. 3.4**), which contains lead SNPs and conditional associations from all our GWAS except automobile speeding propensity (for which it contains a suggestive association: rs141450, $P = 7.88 \times 10^{-8}$). The candidate inversion contains ~50 genes, and 5 of those were significant after Bonferroni correction in the MAGMA gene analysis (**Supplementary Table 12.3**). Among them are *SND1* (Bonferroni-corrected $P = 5.08 \times 10^{-10}$) and *PAX4* (Bonferroni-corrected $P = 1.43 \times 10^{-5}$). *SND1* is implicated as an important factor for normal cell growth⁷³, and *PAX4* is critical to normal fetal development⁷³. The candidate inversion contains genome-wide significant associations in the GWAS Catalog database⁶⁵ with alcohol dependence, educational attainment, and schizophrenia, among other phenotypes.

The fifth notable genomic region is the candidate inversion on chromosome 8 (~7.89 to 11.8 Mb), which contains lead SNPs and conditional associations for all of our GWAS except those of drinks per week and of the first PC of the risky behaviors (**Extended Data Fig. 3.4**). (The strongest associations with drinks per week and the first PC of the risky behaviors have P values of 5.64×10^{-4} and 1.27×10^{-7} , respectively.) There are ~20 genes within the candidate inversion, and interestingly, practically all are significant in the MAGMA gene analysis after Bonferroni-correction (**Supplementary Table 12.3**). Two notable examples are *MSRA* and *CTSB* (with Bonferroni-corrected P values of 2.94×10^{-24} and 4.37×10^{-5} , respectively). *MSRA* is known to be highly expressed in human nervous tissue⁷³, and *CTSB* has a known effect on the processing of an amyloid precursor protein (APP)⁷³. Incomplete processing of APP has been suggested as one of

^p A good example of the strong selection is the differences in allele frequencies across populations for alleles that affect resistance to a lethal form of malaria (see, e.g., ref.²⁶⁹). The protective alleles are very common in areas where the disease is endemic compared to areas where the risk of infection is low or absent.

the causes of Alzheimer's disease⁸³. However, the GWAS Catalog database⁶⁵ does not report any previous associations with Alzheimer's disease within the candidate inversion, or within 500kb of its breakpoints. The GWAS Catalog database⁶⁵ reports genome-wide associations within the breakpoints of the candidate inversion with many other phenotypes, including extraversion, schizophrenia, and chronotype, and the candidate inversion has been analyzed in-depth in a study of neuroticism and depressive symptoms¹⁸.

3.2.2 *Overlap within approximately independent LD blocks*

The observation that several genomic regions contain at least one lead SNP from all or most of our GWAS prompted further investigation. To begin with, we divided the genome into 1703 approximately independent LD blocks identified by Pickrell *et al.*⁸⁴. We then counted the number of blocks that contain lead SNPs from exactly one, two, three, four, five, six, and seven of our main GWAS. Of the 1703 LD blocks, 234 contain at least one lead SNP from exactly one of our seven main GWAS; 93 contain at least one lead SNP from exactly two GWAS; 40 contain at least one lead SNP from exactly three GWAS; 20 contain at least one lead SNP from exactly four GWAS; 7 contain at least one lead SNP from exactly five GWAS; one block (~84.4 Mb to 85.6 Mb on chromosome 3) contains at least one lead SNP from exactly six GWAS; and one block (~51.6 Mb to 55.2 Mb on chromosome 18) contains at least one lead SNP from all seven of our main GWAS. These last two blocks on chromosomes 3 and 18 are respectively located within the first and the second notable genomic region highlighted above and displayed in **Extended Data Fig. 3.4**.

The nine blocks that contain at least one lead SNP from exactly five, six, or seven GWAS each contain at least one general-risk-tolerance lead SNP. Five of these nine blocks overlap with four of the five notable regions described above and are depicted as striped regions in **Extended Data Fig. 3.4**⁹. The five blocks include the two blocks that contain lead SNPs from six and from all seven of our GWAS. The other four blocks that do not overlap with the notable genomic regions are located on chromosome 2 (~44.3 to 46.5 Mb), chromosome 3 (~70.5 to 72.5 Mb), chromosome 6 (~97.8 to 100.6 Mb), and chromosome 7 (113.7 ~ 116.8 Mb). Thus, nine genomic regions contain lead SNPs for at least five of our seven GWAS: the five notable regions described in the previous subsection and the four LD blocks located on chromosomes 2, 3, 6, and 7.

We ran a simulation to benchmark those results and to assess the likelihood of observing such a high level of within-block overlap across the GWAS results, under the null hypothesis that the LD blocks containing the lead SNPs of each GWAS are distributed randomly across the genome and independently from those of the other GWAS. (This null hypothesis is not perfectly realistic: in practice, the phenotypes are phenotypically correlated and their GWAS samples overlap substantially, and some LD blocks are located within regions of the genome that are more likely to contain causal variants. Thus, although informative, this simulation exercise has limitations.)

⁹ There are 27 LD blocks that overlap with the five notable genomic regions, and these are shown, separated by the dotted vertical gray lines, in **Extended Data Fig. 3.4**. Of these 27 blocks, four blocks contain at least one lead SNP from exactly one GWAS; two contain at least one lead SNP from exactly two GWAS; four contain at least one lead SNP from exactly three GWAS; three contain at least one lead SNP from exactly four GWAS; three contain at least one lead SNP from exactly five GWAS; one contains at least one lead SNP from exactly six GWAS; and one contains at least one lead SNP from all seven GWAS (these last two blocks are the ones on chromosomes 3 and 18, located within the first and the second notable genomic regions).

We conducted this analysis with only the general risk tolerance, automobile speeding propensity, drinks per week, ever smoker, and number of sexual partners phenotypes. We excluded adventurousness and the first PC of risky behaviors, because adventurousness is strongly genetically correlated with general risk tolerance, and because the first PC of risky behaviors is constructed from the four risky behaviors. The simulation proceeded in the following way: Since larger LD blocks are more likely to contain lead SNPs for any of our phenotypes, we classified each LD block by length and randomly permuted the LD blocks within each length class independently for each phenotype^r. We then counted the number of LD blocks that contain hits from exactly one, two, three, four, and five GWAS, and we averaged these numbers over 10,000 simulations.

Our results indicate that, under the null hypothesis, we would expect 357.064 LD blocks to contain at least one lead SNP from exactly one GWAS; 43.212 to contain at least one lead SNP from exactly two GWAS; 2.734 to contain at least one lead SNP from exactly three GWAS; 0.077 to contain at least one SNP from exactly four GWAS; and 0.001 to contain at least one lead SNP from all five GWAS. By contrast, the actual number of blocks that contain at least one lead SNP from exactly one, two, three, four, and five of these GWAS are 253, 61, 20, 3, and 1. The expected overlap from our simulation thus differs markedly from the overlap that we actually observe. To investigate this more formally, we conducted a non-parametric Mann-Whitney test⁸⁵ to compare the actual and simulated distributions of within-block overlaps across GWAS. Our results strongly suggest that the distribution of overlap in the simulation is different from the distribution of overlap that we actually observe ($P = 0.0023$).

The simulation exercise thus clearly suggests that the high level of within-block overlap across the results of our seven GWAS are highly unlikely to be due to chance. However, we emphasize again that our null hypothesis is not perfectly realistic and that the simulation exercise has limitations, and that these results should therefore be interpreted with caution.

3.2.3 *Concordance of SNP effects across the nine regions associated with five or more of our phenotypes*

Above, we identified nine genomic regions that contain lead SNPs for at least five of our seven GWAS: the five notable regions on chromosomes 3, 6, 7, 8, and 18, and the four LD blocks located on chromosomes 2, 3, 6, and 7. We investigated whether the signs of the lead SNPs located in these regions tend to be concordant across our primary and supplementary GWAS (in the sense that general-risk-tolerance-increasing alleles are also associated with higher risk taking in the supplementary GWAS, and vice-versa). To do so, we first took the general-risk-tolerance lead SNPs in these nine regions and checked for sign concordance across the six supplementary GWAS. Then, we took the lead SNPs from the six supplementary GWAS in these nine regions and compared their signs to the corresponding signs in the GWAS of general risk tolerance.

These nine regions harbor a total of 37 general-risk-tolerance lead SNPs^s. 217 coefficients of these 37 lead SNPs are available in the results of the six supplementary GWAS (one of the 37 SNPs is

^r We sorted the LD blocks into 9 classes based on the following lengths: 0 to 0.5 Mb (36 blocks), 0.5 to 1 Mb (263 blocks), 1 to 1.5 Mb (526 blocks), 1.5 to 2 Mb (478 blocks), 2 to 2.5 Mb (256 blocks), 2.5 to 3 Mb (82 blocks), 3 to 3.5 Mb (29 blocks), 3.5 to 7.5 Mb (25 blocks), and greater than 7.5 Mb (8 blocks).

^s There are 141 unique lead SNPs across our seven main GWAS in these nine regions. Note that these lead SNPs were obtained using a clumping algorithm with an r^2 threshold of 0.1. See **Supplementary Information sections 2.8** for more details.

only available in the adventurousness GWAS). 205 of these coefficients have concordant signs, 147 are significant at the 5% level in the supplementary GWAS, and 26 are genome-wide significant. Under the null hypothesis that the coefficients are independent across SNPs[†] and have an equal probability of being concordant or discordant across the general risk tolerance and supplementary GWAS, the probability of observing 205 or greater concordant coefficients is very small ($P \leq 9 \times 10^{-47}$). We only found 12 discordant coefficients (9 of which are for drinks per week), and the total number of discordant coefficients is reduced to 4 if for each of the nine regions we exclude coefficients of GWAS that do not contain any lead SNP in the region. None of the discordant coefficients are genome-wide significant, and only 1 is significant at the 5% level (for drinks per week).

The nine regions also harbor 108 SNPs that are lead SNPs for at least one of the six supplementary GWAS. Of these 108 SNPs, some are lead SNPs for more than one of the supplementary GWAS and two are missing in the general-risk-tolerance GWAS, resulting in 111 coefficients that are available for a sign test. 109 of these 111 coefficients are concordant in our GWAS of general risk tolerance. Under the null hypothesis that the coefficients are independent across SNPs have an equal probability of being concordant or discordant across the general risk tolerance and supplementary GWAS, the probability of observing 109 or greater concordant SNPs is very small ($P \leq 3 \times 10^{-30}$). 98 of the concordant coefficients are significant at the 5% level for general risk tolerance, and 31 of these are genome-wide significant; the two discordant coefficients are not significant at the 5% level for general risk tolerance.

Thus, the signs of the lead SNPs located in these regions tend to be highly concordant across our primary and supplementary GWAS, which suggests that these regions represent shared genetic influences on our seven phenotypes (rather than colocalization of causal SNPs).

3.2.4 *Gene overlap across our seven GWAS*

We annotated each lead SNP from our seven GWAS with its most proximate gene, as described in **Supplementary Information section 2.13**. This approach roughly partitions the SNPs throughout the genome into different genomic segments—each associated with a single gene—that can be compared across our main GWAS. When we compared the gene-associated segments identified across our seven GWAS, we found that the only gene segment that is identified in all of them is the one associated with *CADM2* on chromosome 3 (that is located within the long-range LD region on chromosome 3, ~83.4 to 86.9 Mb, that is shared across all our GWAS; **Extended Data Fig. 3.4**).

The second most shared gene segment is the one associated with *TCF4* (discussed in more detail above). *TCF4* is located within the candidate inversion on chromosome 18 (~49.1 to 55.5 Mb) that is shared across all our GWAS (**Extended Data Fig. 3.4**). Lead SNPs from six of our GWAS, but not ever smoker, are annotated to *TCF4*. Three lead SNPs for ever smoker are located within the candidate inversion (of which two are conditional associations) but are instead annotated to the genes *DCC* and *TXNLI*. *DCC* ends ~1.5 Mb before *TCF4*, and *TXNLI* starts ~967 kb after *TCF4*. Four segments, which were associated with the genes *FOXP1*, *MDFIC*, *SIX3*, and *VGLL3* (which

[†] The lead SNPs in any of the nine regions are only approximately independent ($r^2 < 0.1$) from one another, so assuming that the coefficients are independent is an approximation.

is located ~69.8 kb downstream of the notable long-range LD region spanning ~83.4 to 86.9 Mb on chromosome 3), were each identified in five GWAS.

We searched the GWAS Catalog database⁶⁵ for previous associations with the genes associated with these shared segments, and we found some notable associations: *FOXP1* has been associated with ADHD, autism spectrum disorder, chronotype, and schizophrenia; *MDFIC* with BMI and obesity; *SIX3* with fasting plasma glucose, metabolite levels, and myopia; and *VGLL3* with age at menarche and pubertal anthropometrics.

3.3 Results of the discovery GWAS of general risk tolerance

The discovery GWAS of general risk tolerance identified 124 independent genome-wide significant SNPs (i.e., lead SNPs), distributed across chromosomes 1 to 19. The strongest associations were found in the gene *CADM2* on chromosome 3 (rs993137, $P = 2.14 \times 10^{-40}$), on chromosome 7 in the gene *FOXP2* (rs7783012, $P = 7.57 \times 10^{-25}$) and in proximity to the gene *MDFIC* (rs9641536, $P = 5.01 \times 10^{-23}$), and in the gene *MFHAS1* (rs2409095, $P = 1.01 \times 10^{-20}$) on chromosome 8. We report the association results for the 124 lead SNPs in **Supplementary Table 3.1**, together with estimated effect sizes (the β_j 's) expressed in phenotype standard-deviation units per effect-coded allele. **Supplementary Information section 5** reports the results of our successful attempt to replicate these results in our replication GWAS, as well as the results of the “maxFDR” calculation.

As described in **Supplementary Information section 2.8**, in addition to our baseline definition of a lead SNP, we followed Ripke *et al.*⁵⁸ and defined non-overlapping, continuous genomic loci. We identified 99 such loci for general risk tolerance. **Supplementary Table 3.1** lists these loci. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 58,219 SNPs that passed all GWAS quality control filters and that are located within the 99 loci. There were 91 genome-wide significant conditional associations, of which 90 are among the 124 lead SNPs. The single new conditional association, rs163505 (COJO $P = 2.70 \times 10^{-9}$), is genome-wide significant in the discovery GWAS ($P = 1.16 \times 10^{-8}$), but it is in the clump of the lead SNP rs163503. Of the 34 lead SNPs that are not significant in the conditional analysis, 14 are in the notable candidate inversion on chromosome 8 (~7.89 to 11.8 Mb), five are in the notable long-range LD region on chromosome 3 (~83.4 to 86.9 Mb), two are in the notable candidate inversion on chromosome 18 (~49.1 to 55.5 Mb), two are in the notable long-range LD region on chromosome 6 (~25.3 to 33.4 Mb), and two are in other candidate inversion or long-range LD regions. Only 9 of the 34 lead SNPs that are not significant in the conditional analysis are outside long-range LD regions and candidate inversions. Therefore, the exact numbers of lead SNPs within the long-range LD regions and candidate inversions should be interpreted with caution. **Supplementary Table 3.1** reports the results of the conditional analysis.

We display the Manhattan plot of the discovery GWAS in **Fig. 1a** and the quantile-quantile (Q-Q) plot in **Extended Data Fig. 3.2a**. The genomic inflation factor (λ_{GC}) was 1.405 before, and 1.378 after inflation of the standard errors with the square root of the estimated intercept from an LD Score regression ($\widehat{Intercept} = 1.040, SE = 0.011$). As we discuss in **Supplementary Information section 4**, an observed genomic inflation factor larger than 1 is consistent with the expectation that complex traits are polygenic⁸⁶, and an estimated LD Score regression intercept close to unity suggests that the inflation is mainly due to polygenicity rather than to confounding factors⁵³.

All 124 general-risk-tolerance lead SNPs have MAF larger than 1%; only two of the lead SNPs have MAF lower than 5%; four have MAF between 5% and 10%; 33 have MAF between 10% and 25%; and 85 have MAF between 25% and 50%. Thus, the vast majority of our lead SNPs are very common SNPs.

To evaluate the magnitude of the effect-size estimates of the 124 lead SNPs, we compared them with the 124 top associations reported in recent GWAS of height and body mass index (BMI), with the 74 top associations reported in a recent GWAS of educational attainment, and with the 48 top associations reported in a recent GWAS of waist-to-hip ratio adjusted for BMI (WHR)^u. We display the comparison in **Extended Data Fig. 3.3**. The effect-size estimates of the 124 lead SNPs are consistently smaller than the top effect-size estimates of height, BMI, educational attainment, and WHR, both in terms of standard deviation units of the phenotype per effect-increasing allele, as well as in terms of variance explained (R^2). The general-risk-tolerance effect sizes range from 0.008 to 0.026 in phenotype standard-deviation units per effect-increasing allele, compared to a range of 0.038 to 0.191 for height, 0.019 to 0.082 for BMI, 0.014 to 0.048 for educational attainment, and 0.012 to 0.043 for WHR. The variance explained (R^2) by the general-risk-tolerance lead SNPs ranges from 0.003% to 0.019%, compared to 0.036% to 0.283% for height, 0.013% to 0.325% for BMI, 0.010% to 0.036% for educational attainment, and 0.004% to 0.072% for WHR.

We then paired the top SNPs of general risk tolerance with the top SNPs of height, BMI, educational attainment, and WHR (after ranking the SNPs by R^2 for each phenotype), and we calculated the median of the ratio of the R^2 across the paired SNPs. The median ratio was 13.64 between general risk tolerance and height (i.e., for the median ratio, the variance explained was 13.64 times larger for height compared to general risk tolerance), 4.58 between general risk tolerance and BMI, 3.85 between general risk tolerance and WHR, and 2.73 between general risk tolerance and educational attainment. Thus, effect sizes and R^2 of the general-risk-tolerance lead SNPs are substantially lower than those for the comparison phenotypes.

3.3.1 *Genetic correlation between females and males*

We used bivariate LD Score regression⁵³ to calculate the genetic correlation between GWAS performed separately in the sample of females and in the sample of males in the UKB. Our estimate of the genetic correlation ($\hat{r}_g = 0.822$, $SE = 0.033$) is significantly smaller than unity, pointing to some heterogeneity across females and males, but high enough to justify our approach of pooling males and females in our other analyses to maximize statistical power. For further details, see **Supplementary Information section 7.4**.

3.3.2 *Genetic correlations between and heterogeneity of effect sizes across the UKB, 23andMe, and the replication GWAS*

Using bivariate LD Score regression⁵³ we estimated the genetic correlation between the cohort-level GWAS summary statistics from the general-risk-tolerance GWAS in the 23andMe and UKB cohorts, as well as between these GWAS and our replication GWAS (which includes 10 cohorts). Our estimates of the genetic correlations between the 23andMe and UKB GWAS ($\hat{r}_g = 0.767$, SE

^u The data for height, BMI, and WHR are from the publicly available GWAS results of the GIANT consortium, with males and females pooled and restricted to European-ancestry individuals, and the data for educational attainment are from the largest previously published GWAS of educational attainment by Okbay *et al.*¹⁶.

= 0.021), between the 23andMe and the replication GWAS ($\hat{r}_g = 0.759$, $SE = 0.126$), and between the UKB and the replication GWAS ($\hat{r}_g = 0.828$, $SE = 0.135$) are all smaller than unity (though only the first of these estimates is significantly different from unity at the 5% level), pointing to substantial cross-cohort heterogeneity. (For further details, see **Supplementary Information section 7.4**.)

We also used bivariate LD Score regression⁵³ to estimate the genetic correlation between the summary statistics from the discovery GWAS and those from the replication GWAS. We could not reject the null hypothesis of a genetic correlation equal to 1 ($\hat{r}_g = 0.834$, $SE = 0.129$). (For further details, see **Supplementary Information section 7.4**)

Following a suggestion by a Referee and using the methodology described in **Supplementary Information section 2.7**, we evaluated the heterogeneity across the 23andMe and UKB cohorts of the effect-size estimates for the 124 general-risk-tolerance lead SNPs. **Supplementary Table 3.1** reports the P values of Cochran's Q statistic for all 124 lead SNPs, along with the effect-size estimates separately for the 23andMe and UKB cohorts, and **Extended Data Fig. 3.5** plots the effect-size estimates in the 23andMe and UKB cohorts and in the replication GWAS. Cochran's Q statistic is not significant for any of the lead SNPs after Bonferroni-correction for 124 tests (i.e., $P > 0.05/124$ for all 124 lead SNPs). (We note, however, that the power of Cochran's Q test is limited in our setting, because the discovery meta-analysis consists of only two cohorts^{55,56}.) We also generated an omnibus test statistic for heterogeneity by summing the Cochran Q statistics across all lead SNPs⁵⁷. As mentioned in **Supplementary Information section 2.7**, the sum of the Q statistics of the 124 lead SNPs is (approximately) χ^2 -distributed with 124 degrees of freedom. We obtained a sum of 195.64, with a corresponding P value of 4.32×10^{-5} (under the null hypothesis of homogeneity, the expected value of that sum is equal to the number of degrees of freedom, which here is 124). Consistent with our genetic correlation estimate of less than unity between the 23andMe and UKB cohorts, this points to the presence of some heterogeneity across the 23andMe and UKB cohorts.

3.3.3 *Results of the cross-lookup of the general-risk-tolerance lead SNPs and loci in the supplementary GWAS*

The results of the cross-lookup (described in **Supplementary Information section 2.12**) suggest substantial genetic overlap between the lead SNPs of our primary GWAS and those of our supplementary GWAS. 72 of the 124 general-risk-tolerance lead SNPs are also lead SNPs, or in LD ($r^2 > 0.1$) with a lead SNP, for at least one of the other main phenotypes, and 46 of the 99 general-risk-tolerance loci contain a lead SNP for at least one of the other main phenotypes. The overlap is particularly large between general risk tolerance and adventurousness: 45 of the 124 general-risk-tolerance lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for adventurousness. We also found that 49 of the 124 general-risk-tolerance lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), of one of the four risky behaviors or their first PC.

To benchmark the likelihood of observing such substantial genetic overlap between the lead SNPs and loci of our primary and supplementary GWAS, we conducted a resampling exercise under the null hypothesis that the lead SNPs of our supplementary GWAS are distributed randomly across the genome and independently from the general-risk-tolerance lead SNPs and loci and from the lead SNPs of the other GWAS. The resampling exercise involved 10,000 runs. In each run, for each lead SNP of the supplementary GWAS we randomly selected an autosomal SNP matched on

MAF (with five-percentage-point MAF windows). We then counted how many of the 124 general-risk-tolerance lead SNPs were in weak LD ($r^2 > 0.1$) with at least one of the resampled lead SNPs, as well as how many of the 99 general-risk-tolerance loci contained at least one of the resampled lead SNPs, and took the average count across the 10,000 runs. Our results imply that, under the null hypothesis, we would expect 10.3 of the 124 general-risk-tolerance lead SNPs to be in weak LD with a lead SNP for at least one of the supplementary phenotypes; 2.2 to also be in weak LD with a lead SNP for adventurousness; and 8.1 to also be in weak LD with a lead SNP for at least one of the four risky behaviors or their first PC. Similarly, under the null hypothesis, we would expect 3.5 of the 99 general-risk-tolerance loci to contain a lead SNP for at least one of the supplementary phenotypes. In all four cases, we strongly reject the null hypothesis ($P < 0.0001$ in all four cases). The overlap we observe between the 124 general-risk-tolerance lead SNPs and the lead SNPs from the other GWAS is thus much greater than what could be expected by chance.

3.3.4 *Novelty of the lead SNPs of the discovery GWAS of general risk tolerance*

We assessed the novelty of the lead SNPs by performing a lookup in the NHGRI-EBI GWAS Catalog database⁶⁵ for each of the 124 general-risk-tolerance lead SNPs (and the SNPs in LD, $r^2 > 0.1$), searching specifically for previous associations with general risk tolerance (and similar phenotypes) (see **Supplementary Information section 2.10** for details). We found that the GWAS Catalog database contained no previous associations with general risk tolerance (or similar phenotypes).

However, we know of one previous study by Day *et al.*⁶⁶ that analyzed the same general-risk-tolerance phenotype measure (which they refer to as “risk-taking propensity”) as the one we study, in the first release of the UKB data. Day *et al.* report one independent lead SNP (rs4856591) associated with general risk tolerance in the gene *CADM2*. This association was replicated by Boutwell *et al.*⁶⁸ using a proxy SNP available in an independent sample (rs1865251) located ~125 kb from the original lead SNP (i.e., rs4856591).

The original lead SNP from ref.⁶⁶ is not available in our main reference panel (and therefore not available in the summary statistics from our discovery GWAS), but the proxy SNP from ref.⁶⁸ is available. Our discovery GWAS also contains a proximate SNP (rs4856590) only 22 bp away from the original lead SNP from ref.⁶⁶. Both rs1865251 and rs4856590 are genome-wide significant in our discovery GWAS. They are “clumped” with our top lead SNP rs993137 ($P = 2.14 \times 10^{-40}$), and the LD (r^2) between our lead SNP rs993137 and rs1865251 and rs4856590 is 0.996 and 0.949, respectively. Because its locus is associated with general risk tolerance in a previous peer-reviewed publication, we do not consider our lead SNP rs993137 to be a novel association.

We also know of one concurrent GWAS by Strawbridge *et al.*⁶⁷ on general risk tolerance (which Strawbridge *et al.* refers to as “risk-taking behavior”). Strawbridge *et al.* also analyzed the same phenotype measure as the one we and Day *et al.*⁶⁶ study, but used a somewhat different set of individuals than Day *et al.* in the first release of the UKB data. They identified one independent lead SNP on chromosome 6 (i.e., rs9379971) in addition to the previous association in *CADM2* from ref.⁶⁶ and ref.⁶⁸. The SNP rs9379971 is genome-wide significant in our discovery GWAS, and it is “clumped” with a lead SNP of the discovery GWAS (rs1417998, $P = 2.92 \times 10^{-10}$).

Thus, to the best of our knowledge, all of the loci we identified through our GWAS of general risk tolerance are novel associations with general risk tolerance, except for our top lead SNP rs993137^v. We therefore consider 123 of the 124 lead SNPs to be newly identified associations with general risk tolerance (and similar phenotypes). We report the novelty of our 124 lead SNPs in the column “New association” in **Supplementary Table 3.1**.

3.3.5 Results of the GWAS Catalog lookup of the lead SNPs of the primary GWAS of general risk tolerance

To investigate the potential overlap between general risk tolerance and other phenotypes, we performed a lookup for each of our 124 lead SNPs (and the SNPs in LD, $r^2 > 0.6$) in the NHGRI-EBI GWAS Catalog database⁶⁵ (see **Supplementary Information section 2.11** for details). The results are reported in **Supplementary Table 3.4**. In total, we found 61 overlaps between our lead SNPs (and the SNPs in LD, $r^2 > 0.6$) and the previous associations reported in the GWAS Catalog database. Some of our lead SNPs overlap with multiple previous associations: the 61 overlaps involve only 27 of our 124 lead SNPs.

Notably, we found overlaps at five schizophrenia loci, one bipolar disorder locus, four educational attainment loci, and two loci associated with cognitive function and information processing speed. The multiple overlaps with schizophrenia are consistent with the 16 second-stage hits we obtained for schizophrenia in the proxy-phenotype analysis (**Supplementary Table 8.1**), as well as with our finding of strong joint enrichment for association with schizophrenia ($P = 3.0 \times 10^{-9}$) among our general-risk-tolerance lead SNPs on a non-parametric Mann-Whitney test (**Supplementary Information section 8**). Further, 70% of the 122 general-risk-tolerance lead SNPs available in the schizophrenia summary statistics have concordant signs for the two phenotypes ($P = 8.3 \times 10^{-6}$). (We also note that general risk tolerance and schizophrenia are moderately and positively genetically correlated ($\hat{r}_g = 0.173$, $SE = 0.021$; **Supplementary Information section 7**)). However, we caution that it is possible that this overlap with the schizophrenia results is primarily attributable to the fact that the schizophrenia GWAS⁵⁸ was relatively well-powered, rather than to shared genetic etiology between risk tolerance and schizophrenia.

In addition, we found overlaps at one ADHD locus, one extraversion locus, and one brain volume (superior frontal gyrus grey matter volume) locus. Other potentially interesting overlaps were found at a resting heart rate locus and a motion sickness locus. Several traits associated with autoimmune disease also overlap with our lead SNPs: two loci associated with cholangitis, one locus associated with ulcerative colitis/atopic dermatitis, and one locus associated with type 1 diabetes (that locus is also associated with height, age at menarche, and male pattern baldness). We also found overlaps at one locus associated with age at menarche, at two loci associated with carcinoma, and at one locus associated with breast cancer.

The remaining overlaps are with seemingly unrelated traits, for example at loci associated with tooth development (one locus), gut microbiota diversity (one locus), blood protein levels (one locus), blood pressure and type 2 diabetes (one locus), and ear infection (one locus). The genetic overlap of general risk tolerance with a large number of traits underlines the possibility of widespread pleiotropy in the human genome.

^v Since we consider the GWAS by Strawbridge *et al.*⁶⁷ to be concurrent, we consider our lead SNP rs1417998 to be a novel association with general risk tolerance.

We note that both the existence and absence of overlaps between any two phenotypes should be interpreted with care. The existence of overlaps is not necessarily evidence of a shared genetic architecture. For example, a shared tagging variant could be in LD with two different causal variants that each only cause one of the traits⁸⁷. Further, the absence of overlaps may be the result of inadequate statistical power in the currently available GWAS for either phenotype.

3.3.6 *General-risk-tolerance lead SNPs in long-range LD regions*

8 of the 15 long-range LD regions identified by Price *et al.*⁶⁰ (**Supplementary Information section 2.9**) contain lead SNPs for at least one of our primary or supplementary GWAS (of general risk tolerance, adventurousness, the four risky behaviors, and their first PC). We focus this section on the two long-range LD regions that contain general-risk-tolerance lead SNPs and that are notable because they also contain lead SNPs for all or most of our other six main GWAS phenotypes: the long-range LD regions ~83.4 to 86.9 Mb on chromosome 3 (**Extended Data Fig. 3.4**) and ~25.3 to 33.4 Mb on chromosome 6 (**Extended Data Fig. 3.4**). These two regions are both among the five notable genomic regions we highlighted in **Supplementary Information section 3.2**. Those two regions contain 10 of the 124 general-risk-tolerance lead SNPs. We also report a third long-range LD region (~135.5 to 138.8 Mb on chromosome 5) that contains two general-risk-tolerance lead SNPs (but no lead SNPs for the other phenotypes), and below in **Supplementary Information section 3.4** we present the five remaining long-range LD regions that we identified in our other six main GWAS. Only one long-range LD region (i.e., chromosome 3, ~83.4 to 86.9 Mb, **Extended Data Fig. 3.4**) contains lead SNPs for all seven of our phenotypes. We note again that the exact numbers of lead SNPs within the long-range LD regions should be interpreted with caution because many of the lead SNPs within these regions are not conditional associations.

The first notable long-range LD region, spanning ~83.4 to 86.9 Mb on chromosome 3, is displayed in a local Manhattan plot in **Extended Data Fig. 3.4** and was discussed in **Supplementary Information section 3.2**. The region is noteworthy because it contains more than one lead SNP for all of our GWAS, and it contains six lead SNPs for general risk tolerance, of which three are located within the *CADM2* gene (~85.0 to 86.2 Mb), two are located ~76 kb and ~115 kb upstream of *CADM2*, and one is located ~547 kb downstream of *CADM2*, closer to the gene *VGLL3*. (The gene *VGLL3*, ~86.9 to 87.0 Mb on chromosome 3, is ~69.8 kb downstream of the long-range LD region.) As we discussed in **Supplementary Information section 3.2**, the long-range LD region contains only one gene—*CADM2*, which covers ~1.2 Mb of the ~3.5 Mb long-range LD region. *CADM2* was the most significantly associated gene in the MAGMA gene analysis (Bonferroni-corrected $P = 1.09 \times 10^{-50}$, **Supplementary Table 12.3**), and it contains our strongest association with general risk tolerance (rs993137, $P = 2.14 \times 10^{-40}$). The GWAS Catalog database⁶⁵ reports many genome-wide significant associations within the long-range LD region, including age at menarche, BMI, educational attainment, and information processing speed. Most of the previous associations within the long-range LD region are annotated to *CADM2*, which encodes a synaptic cell adhesion molecule⁷³, and is related to synapse formation⁷⁴ and brain plasticity⁷⁵.

The second long-range LD region (~25.3 to 33.4 Mb on chromosome 6) covers all the Human Leukocyte Antigen (HLA) genes⁷³. It was also discussed in **Supplementary Information section 3.2**, and it is displayed in a local Manhattan plot in **Extended Data Fig. 3.4**. It contains four lead SNPs located in (or in close proximity to) four different genes: *HIST1H2AC*, *HIST1H2BD*, *TRIM27*, and *C4B*, of which the first three are significant in the MAGMA gene analysis after

Bonferroni-correction (**Supplementary Table 12.3**). The region is notable because it contains lead SNPs for all our other main GWAS, except drinks per week for which we identified a suggestively associated SNP (rs6937318, $P = 3.83 \times 10^{-7}$). As we discussed in **Supplementary Information section 3.2**, the region contains ~250 genes, and the MAGMA gene analysis found ~30 significant genes in the region after Bonferroni-correction (however, none of the actual HLA genes are significant, **Supplementary Table 12.3**). The GWAS Catalog database⁶⁵ reports more than a thousand genome-wide significant associations across hundreds of traits within the long-range LD region (~25.3 to 33.4 Mb on chromosome 6), including Alzheimer's disease, autism spectrum disorder, educational attainment, and schizophrenia.

We also identified a third long-range LD region (~135.5 to 138.8 Mb on chromosome 5) that contains two lead SNPs associated with general risk tolerance within the genes *CTNNA1* and *ETFL*, but the region does not contain lead SNPs for any of our other main GWAS.

3.3.7 *General-risk-tolerance lead SNPs in candidate inversions*

Of the 154 genomic segments deemed highly prone to inversion polymorphisms (i.e., the 154 “candidate inversions,” described in **Supplementary Information section 2.9**), we identified 44 that contain lead SNPs for at least one of our GWAS. 13 of these 44 candidate inversions contain a total of 30 general-risk-tolerance lead SNPs. We note again that the exact numbers of lead SNPs within the candidate inversions should be interpreted with caution because many lead SNPs within those regions are not conditional associations.

This section focuses on four of the 13 candidate inversions that we find noteworthy because they contain lead SNPs for general risk tolerance as well as for most of our other main GWAS. Three of these four candidate inversions are among the five notable genomic regions we highlighted in **Supplementary Information section 3.2**. (Below, in **Supplementary Information section 3.4**, we discuss three other of the 44 candidate inversions of the 44, which are notable because they are shared across four of our GWAS, excluding general risk tolerance.) The four candidate inversions that we discuss in this section span ~124.6 to 132.7 Mb on chromosome 7 (**Extended Data Fig. 3.4**), ~7.89 to 11.8 Mb on chromosome 8 (**Extended Data Fig. 3.4**), ~70.1 to 74.4 Mb on chromosome 16, and ~49.1 to 55.5 Mb on chromosome 18 (**Extended Data Fig. 3.4**). Only one of these candidate inversions (i.e., ~49.1 to 55.5 Mb on chromosome 18, **Extended Data Fig. 3.4**) contains lead SNPs and conditional associations for all our GWAS.

The first candidate inversion is on chromosome 7 (~124.6 to 132.7 Mb). We discussed it in **Supplementary Information section 3.2** and we display it in a local Manhattan plot in **Extended Data Fig. 3.4**. This candidate inversion is notable because it contains lead SNPs and conditional associations for all our GWAS, except automobile speeding propensity (for which the strongest association is almost genome-wide significant (rs141450, $P = 7.88 \times 10^{-8}$)). The candidate inversion contains one lead SNP for general risk tolerance, two lead SNPs for adventurousness, one for drinks per week, one for ever smoker, one for number of sexual partners, and one for the first PC of the risky behaviors. As we discussed in **Supplementary Information section 3.2**, the candidate inversion contains ~50 genes, and five of those were significant after Bonferroni correction in the MAGMA gene analysis (**Supplementary Table 12.3**). The GWAS Catalog database⁶⁵ reports previous associations with traits such as alcohol dependence, educational attainment, and schizophrenia.

The second candidate inversion spans ~7.89 to 11.79 Mb on chromosome 8. We discussed it in **Supplementary Information section 3.2** and we display it in a local Manhattan plot in **Extended Data Fig. 3.4**. This candidate inversion is notable, because it contains lead SNPs and conditional associations for all our GWAS, except drinks per week and the first PC of the risky behaviors. (The strongest association with drinks per week within the candidate inversion is rs574968044, $P = 5.64 \times 10^{-4}$, and the strongest association with the first PC of the risky behaviors is rs2898249, $P = 1.27 \times 10^{-7}$.) The candidate inversion contains 15 lead SNPs for general risk tolerance, five for adventurousness, one for automobile speeding propensity, two for ever smoker, and four for number of sexual partners. As we discussed in **Supplementary Information section 3.2**, the candidate inversion contains ~20 genes, of which practically all were significant after Bonferroni correction in the MAGMA gene analysis (**Supplementary Table 12.3**). The GWAS Catalog database⁶⁵ reports genome-wide associations in the candidate inversion with many behavioral phenotypes, including extraversion, neuroticism, schizophrenia, and chronotype.

The third candidate inversion is on chromosome 18 (~49.1 to 55.5 Mb) and was also discussed in **Supplementary Information section 3.2**. It contains lead SNPs and conditional associations for all our GWAS, and it contains more than one lead SNP for all our GWAS except drinks per week, for which it contains only one. We display the region in a local Manhattan plot in **Extended Data Fig. 3.4**. The candidate inversion contains three lead SNPs for general risk tolerance, two for adventurousness, two for automobile speeding propensity, one for drinks per week, three for ever smoker, eight for number of sexual partners, and four for the first PC of the risky behaviors. As we discussed in **Supplementary Information section 3.2**, among the ~20 genes within the candidate inversion, we find only one gene—*TCF4* (Bonferroni-corrected $P = 5.51 \times 10^{-9}$)—significant after Bonferroni correction in the MAGMA gene analysis (**Supplementary Table 12.3**). *TCF4* plays an important role in nervous system development, and mutations in the gene are known to cause the rare Pitt-Hopkins syndrome⁷³. The syndrome is characterized by distinct facial features, intellectual disability, delayed motor skills, and epilepsy, among many other symptoms^{77–79}. The GWAS Catalog database⁶⁵ reports genome-wide significant associations mapped to *TCF4* with schizophrenia and reports previous genome-wide associations within the candidate inversion with traits such as autism spectrum disorder, ADHD, depression, educational attainment, schizophrenia, and subcortical brain region volumes.

A fourth candidate inversion is on chromosome 16 (~70.1 to 74.4 Mb) and contains lead SNPs and conditional associations for the GWAS of general risk tolerance, ever smoker, number of sexual partners, and the first PC of the risky behaviors. (The strongest association with adventurousness within the candidate inversion is rs9929242 ($P = 2.21 \times 10^{-6}$), the strongest association with automobile speeding propensity is rs2158268 ($P = 3.35 \times 10^{-6}$), and the strongest association with drinks per week is rs11648570 ($P = 1.50 \times 10^{-7}$.) The candidate inversion contains ~35 genes, of which 2 are significant after Bonferroni correction in the MAGMA gene analysis (**Supplementary Table 12.3**): *CHST4* (Bonferroni-corrected $P = 4.69 \times 10^{-3}$) and *CMTR2* (Bonferroni-corrected $P = 0.024$). *CHST4* is involved in normal cell function via carbohydrate sulfotransferase⁷³, and *CMTR2* is involved in methyltransferase⁷³. There is also one additional gene ~458 kb upstream of the candidate inversion—*NFAT5*—that is significant after Bonferroni correction in the MAGMA gene analysis (Bonferroni-corrected $P = 4.66 \times 10^{-3}$). It is notable because it plays a central role in gene transcription during immune response⁷³, consistent with the significant estimate of the category “Immune/Hematopoietic” in the partitioning of the SNP heritability into functional categories with LD Score regression (**Supplementary Information section 12**). The candidate

inversion contains genome-wide significant associations in the GWAS Catalog database⁶⁵ with total cholesterol, prostate cancer, and stroke, among other phenotypes.

The remaining nine of the 13 candidate inversions that contain general-risk-tolerance lead SNPs each contain only one general-risk-tolerance lead SNP, and the overlap with the other GWAS is low.

3.3.8 General-risk-tolerance lead SNPs in LD with 1000 Genomes structural variants

In addition, 24 of the 124 lead SNPs are located within, or in strong LD with another variant within, 23 different 1000G structural variants. All candidate inversions that contain general-risk-tolerance lead SNPs, as well as these 1000G structural variants, are reported in **Supplementary Table 3.1**.

3.3.9 Interaction of age and sex with general-risk-tolerance lead SNPs

Following a suggestion by a Referee, we investigated whether there is evidence of interaction effects between each of the 124 general-risk-tolerance lead SNPs and either age or sex. We began by assessing the statistical power to detect such interaction effects, with a series of power calculations. We assumed that the interaction effects could have R^2 's that are 5%, 10%, 25%, 50%, or 100% as large as the largest and smallest R^2 's of the 124 general-risk-tolerance lead SNPs. The power to find an interaction effect at the 5% level of significance was estimated to be 53.8%, 82.8%, 99.6%, 100%, and 100%, respectively, for the largest effect size ($R^2 \sim 0.02\%$), and 13.2%, 22.1%, 47.0%, 76.0%, 96.5%, respectively, for the smallest effect size ($R^2 \sim 0.003\%$). If we account for multiple hypothesis testing and instead use a significance level of 5%/124, the power ranges from 6.9% to 100% for the largest effect size, and from 0.4% to 59% for the smallest. The statistical power to detect an interaction effect is thus not very high, but may still be sufficient, depending on the size of the interaction effect.

For each of the 124 lead SNPs, we performed two regressions: (1) one regression of general risk tolerance on the SNP and the interaction between the SNP and sex; and (2) one regression of general risk tolerance on the SNP and the interactions with two of three age bins (to test for nonlinear age effects, we used three age bins: ≤ 50 , 51–60, and ≥ 61). Both regressions also included our standard set of covariates (described in **Supplementary Table 2.2**). For none of the 124 lead SNPs was the interaction with sex statistically significant after Bonferroni correction for 124 tests. Similarly, for none of the lead SNPs were the interactions with the two age bins nominally significant after correction for $124 \times 2 = 248$ tests.

3.4 Results of the supplementary GWAS

Our six supplementary GWAS—of adventurousness, the four risky behaviors, and of the first PC of the four risky behaviors—identified a total of 740 lead SNPs^w. We consider 729 of these 740 lead SNPs to be novel associations for these phenotypes. The association results are reported in **Supplementary Table 3.2**. We identified 167 lead SNPs for adventurousness, 42 for automobile

^w As mentioned above, we did not attempt replication of the results of our six supplementary GWAS in independent data, because we did not have access to such data for the six supplementary phenotypes. However, as we report in **Supplementary Information section 5**, we calculated the “maxFDR”⁷¹, an upper bound on the false discovery rate (FDR), for each GWAS. The maxFDR estimates were low across all GWAS, thus providing reassurance about the robustness of the lead associations.

speeding propensity, 85 for drinks per week, 223 for ever smoker, 117 for number of sexual partners^x, and 106 for the first PC of the risky behaviors.

The results are displayed in Manhattan plots in **Extended Data Fig. 3.1** and in Q-Q plots in **Extended Data Fig. 3.2**. The genomic inflation factors (λ_{GC}) across the phenotypes ranges from 1.254 to 1.470, and the estimated LD Score regression intercepts were in the range of 1.026 to 1.051, consistent with polygenicity and low levels of confounding from population stratification^{53,86}. The estimated effect sizes (the $\hat{\beta}_j$'s) reported in **Supplementary Table 3.2** are expressed in phenotype standard deviation units per effect-coded allele. (The phenotype standard deviations are reported in **Supplementary Table 1.2**).

3.4.1 GWAS of adventurousness

We identified 167 lead SNPs associated with adventurousness. The strongest associations were found within, or in close proximity to, the genes *CADM2* (rs10433500, $P = 9.31 \times 10^{-84}$), *SATB1* (rs13090941, $P = 3.26 \times 10^{-21}$), and *FOXP2* (rs10228494, $P = 1.23 \times 10^{-19}$). 47 of the 167 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for general risk tolerance. 52 of the 167 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for the four risky behaviors, and their first PC.

To the best of our knowledge, there are no previous studies with genome-wide significant associations for adventurousness, and we consider all of our lead SNPs to be novel associations. We report the novelty of our adventurousness lead SNPs (and of the lead SNPs of the other supplementary GWAS) in the column “New locus” in **Supplementary Table 3.2**.

We identified 137 loci (**Supplementary Information section 2.8**) for adventurousness. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 69,804 SNPs that passed all GWAS quality control filters and that are located within the 137 loci. There were 126 genome-wide significant conditional associations, of which 120 are among the 167 lead SNPs. The six new conditional associations (COJO $P = 8.69 \times 10^{-10}$ – 3.36×10^{-8}), are all genome-wide significant in the GWAS ($P = 1.73 \times 10^{-12}$ – 3.26×10^{-8}), but they are in the clumps of other lead SNPs. **Supplementary Table 3.2** lists the loci and reports the results of the conditional analysis.

We now discuss long-range LD and candidate inversion regions that contain adventurousness lead SNPs. We emphasize, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations. 18 of the 167 lead SNPs are distributed across three of the 15 long-range LD regions identified by Price *et al.*⁶⁰, of which two are the long-range LD regions that we describe above in **Supplementary Information section 3.2** on chromosome 3 (~83.4 to 86.9 Mb) and chromosome 6 (~25.3 to 33.4 Mb). Remarkably, 16 of these 18 lead SNPs are located within the long-range LD region on chromosome 3. Most of these 16 SNPs are located within, or in proximity to, the *CADM2* gene, but three of them are closer to the gene *VGLL3* (which is located ~69.8 kb downstream of the long-range LD region). The third long-range LD region, on chromosome 8 (~111.9 to 114.9 Mb), does not contain lead SNPs for any of our other six main GWAS. There is also a total of 24 lead SNPs located within 16 candidate inversions.

^x As mentioned in a previous footnote, we excluded one of the 118 number-of-sexual-partners lead SNPs we initially identified (see **Supplementary Information section 3.4.5** for details).

Furthermore, there are 29 lead SNPs located within, or in strong LD with another variant within, 29 different 1000G structural variants. All candidate inversions that contain adventurousness lead SNPs, as well as these 1000G structural variants, are reported in **Supplementary Table 3.2**.

3.4.2 *GWAS of automobile speeding propensity*

We identified 42 lead SNPs associated with automobile speeding propensity. The strongest association was found within the gene *CADM2* (rs17516256, $P = 9.9 \times 10^{-21}$). Seven of the 42 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for general risk tolerance. 17 of the 42 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for at least one of the other main phenotypes we analyze, excluding the first PC of the risky behaviors.

To the best of our knowledge, there are no previous studies with genome-wide significant associations for automobile speeding propensity, so we consider all of our lead SNPs to be novel associations (**Supplementary Table 3.2**).

We identified 36 loci (**Supplementary Information section 2.8**) for automobile speeding propensity. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 38,156 SNPs that passed all GWAS quality control filters and that are located within the 36 loci. There were 33 genome-wide significant conditional associations, of which 30 are among the 42 lead SNPs. The three new conditional associations (COJO $P = 3.32 \times 10^{-10}$ – 2.48×10^{-8}), are all genome-wide significant in the GWAS ($P = 3.30 \times 10^{-12}$ – 3.55×10^{-8}), but they are in the clumps of other lead SNPs. **Supplementary Table 3.2** lists the loci and reports the results of the conditional analysis.

We now discuss long-range LD and candidate inversion regions that contain automobile-speeding-propensity lead SNPs. We emphasize again, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations. Eight of the 42 lead SNPs are distributed across two of the long-range LD regions identified by Price *et al.*⁶⁰. Those two regions are the two long-range LD regions we described above in **Supplementary Information section 3.2** and that are shared across general risk tolerance and all or most of our GWAS. Three of the eight lead SNPs are in the region on chromosomes 3 (~83.4 to 86.9 Mb), and the other five are in the region on chromosome 6 (~25.3 to 33.4 Mb). The three lead SNPs located in the long-range LD region on chromosome 3 (~83.4 to 86.9 Mb) are all within the *CADM2* gene. There is also a total of 24 lead SNPs located within 16 candidate inversions. Furthermore, seven lead SNPs are located within, or in strong LD with another variant within, seven different 1000G structural variants. All candidate inversions that contain automobile speeding propensity lead SNPs, as well as these 1000G structural variants, are reported in **Supplementary Table 3.2**.

3.4.3 *GWAS of drinks per week*

We identified 85 lead SNPs associated with drinks per week. The strongest and most notable of these associations is located within the alcohol dehydrogenase 1B gene on chromosome 4 (*ADH1B*, rs1229984, $P = 7.8 \times 10^{-202}$). Multiple genome-wide significant associations were also found within, or in close proximity to, other alcohol dehydrogenase genes—*ADH1A* (rs62307263, $P = 3.42 \times 10^{-11}$), *ADH1C* (rs113659074, $P = 9.93 \times 10^{-23}$), and *ADH7* (rs114112910, $P = 1.08 \times 10^{-8}$). Four of the 85 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for general

risk tolerance. 21 of the 85 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for at least one of the other main phenotypes we analyze, excluding the first PC of the risky behaviors.

Following the procedure described in **Supplementary Information section 2.10**, we found that the lead SNPs located within, or in close proximity to, the genes *ADH1A*, *ADH1B*, *ADH1C*, *ADH7*, *GCKR*, and *KLB*, have previously been associated with alcohol consumption (or similar phenotypes), and we therefore do not consider associations annotated to these genes to be novel associations with drinks per week (**Supplementary Table 3.2**).

We identified 62 loci (**Supplementary Information section 2.8**) for drinks per week. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 111,146 SNPs that passed all GWAS quality control filters and that are located within the 62 loci. There were 61 genome-wide significant conditional associations, of which 56 are among the 85 lead SNPs. The five new conditional associations (COJO $P = 5.19 \times 10^{-28}$ – 2.09×10^{-8}), are all genome-wide significant in the GWAS ($P = 7.42 \times 10^{-24}$ – 2.07×10^{-8}), but they are in the clumps of other lead SNPs. **Supplementary Table 3.2** lists the loci and reports the results of the conditional analysis.

We now discuss long-range LD and candidate inversion regions that contain drinks-per-week lead SNPs. We emphasize again, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations. Two of the 85 lead SNPs are located within the same long-range LD region on chromosome 3 (~83.4 to 86.9 Mb) that contains lead SNPs for all our GWAS phenotypes (**Supplementary Information section 3.2**), and both lead SNPs are located within *CADM2*. Unlike for the six other main phenotypes we analyze, there are no lead SNPs for drinks per week in the long-range LD region spanning ~25.3 to 33.4 Mb on chromosome 6, but the region contains a suggestive association at rs6937318 (P value = 3.83×10^{-7}). However, as can be seen in **Extended Data Fig. 3.4**, the general level of association is much lower for drinks per week in that long-range LD region in comparison with the other main phenotypes we analyze.

15 lead SNPs are located across 15 candidate inversions. Of these, none is located within the candidate inversion spanning ~7.89 to 11.8 Mb on chromosome 8, unlike for general risk tolerance, adventurousness, ever smoker, and number of sexual partners. (The strongest association with drinks per week within that candidate inversion is rs574968044, ($P = 5.64 \times 10^{-4}$).) Just as for the long-range LD region on chromosome 6 (~25.3 to 33.4 Mb), the general level of association is much lower for drinks per week in the candidate inversion on chromosome 8 in comparison with the other main phenotypes, as can be seen in **Extended Data Fig. 3.4**.

15 lead SNPs are located within, or in strong LD with another variant within, 20 different 1000G structural variants (some lead SNPs are in strong LD with SNPs in multiple structural variants). All candidate inversions that contain drinks per week lead SNPs, as well as these 1000G structural variants, are reported in **Supplementary Table 3.2**.

3.4.4 GWAS of ever smoker

We identified 223 lead SNPs associated with ever smoker. The strongest associations are located within, or in close proximity to, the genes *NCAMI* (rs7938812, $P = 7.1 \times 10^{-48}$), *ZEB2* (rs961414, $P = 6.99 \times 10^{-28}$), *REV3L* (rs240955, $P = 3.84 \times 10^{-23}$), *NT5C2* (rs7092200, $P = 7.44 \times 10^{-22}$), *CLU* (rs11783093, $P = 7.44 \times 10^{-22}$), *TMEM182* (rs1368550, $P = 1.08 \times 10^{-21}$), and *CADM2* (rs34495106, $P = 2.23 \times 10^{-20}$). There are three additional lead SNPs located within, or in close proximity to,

CADM2. 22 of the 223 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for general risk tolerance. 59 of the 223 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for at least one of the other main phenotypes we analyze, excluding the first PC of the risky behaviors.

Following the procedure described in **Supplementary Information section 2.10**, we found that the two lead SNPs located within the genes *NCAM1* and *BDNF* have previously been associated with smoking behavior. We therefore do not consider those two loci to be novel associations with ever smoker (**Supplementary Table 3.2**).

Previous GWAS on nicotine dependence⁸⁸ and number of cigarettes per day (CPD)^{35,88} report genes in the nicotine receptor gene family (i.e., the *CHRN* family³⁵) as some of their strongest associations. In our results, none of the nicotine receptor genes (*CHRN*) contained any lead SNPs, consistent with the theory that the ever smoker phenotype is more strongly mediated by risk tolerance and social influences than by vulnerability to nicotine addiction⁸⁹.

We identified 183 loci (**Supplementary Information section 2.8**) for ever smoker. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 133,268 SNPs that passed all GWAS quality control filters and that are located within the 183 loci. There were 172 genome-wide significant conditional associations, of which 162 are among the 223 lead SNPs. The ten new conditional associations (COJO $P = 5.19 \times 10^{-28}$ – 2.09×10^{-8}), are all genome-wide significant in the GWAS (3.50×10^{-14} – 4.21×10^{-8}), but they are in the clumps of other lead SNPs. **Supplementary Table 3.2** lists the loci and reports the results of the conditional analysis.

We now discuss long-range LD and candidate inversion regions that contain ever-smoker lead SNPs. We emphasize again, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations. 14 lead SNPs are located in six of the long-range LD regions identified by Price *et al.*⁶⁰. These include the long-range LD regions ~83.4 to 86.9 Mb on chromosome 3 and ~25.3 to 33.4 Mb on chromosome 6, which were also found to contain lead SNPs for general risk tolerance and most of our other GWAS phenotypes (**Supplementary Information section 3.2**). The four remaining long-range LD regions are on chromosome 1 (~48.2 to 52.2 Mb), chromosome 2 (~134.7 to 138.2 Mb), chromosome 12 (~111.0 to 113.5 Mb), and chromosome 20 (~32.5 to 35.0 Mb). 24 lead SNPs are located within 16 candidate inversions. There are also 56 lead SNPs located within, or in strong LD with another variant within, 69 different 1000G structural variants (some lead SNPs are in strong LD with SNPs in multiple structural variants). All candidate inversions that contain ever smoker lead SNPs, as well as these 1000G structural variants, are reported in **Supplementary Table 3.2**.

We also performed a replication of the eight genome-wide significant SNPs from the GWAS of ever smoker by the TAG consortium³⁵ in our GWAS in the UKB only (and not in our meta-analyses of our UKB GWAS with the TAG summary statistics, as our replication sample must be independent of the TAG sample). All eight SNPs have concordant signs, and all are highly significant. Specifically, six of the eight SNPs are genome-wide significant (P value $< 5 \times 10^{-8}$) in our GWAS of ever smoker in the UKB (rs6265, $P = 1.1 \times 10^{-14}$; rs4923457, $P = 2.3 \times 10^{-11}$; rs4923460, $P = 1.8 \times 10^{-11}$; rs4074134, $P = 2.6 \times 10^{-11}$; rs6484320, $P = 2.1 \times 10^{-11}$; rs879048, $P = 1.2 \times 10^{-11}$), and the two remaining SNPs are suggestively significant (rs1013442, $P = 1.5 \times 10^{-7}$; rs1304100, $P = 1.2 \times 10^{-7}$). It should be noted that the eight SNPs reported in the TAG consortium

results are not independent, and if we apply our definition of lead SNPs from **Supplementary Information section 2.8**, then the SNP rs6265 would be the only lead SNP, and the other seven SNPs would be clumped to that SNP. rs6265 is one of the lead SNPs in our GWAS of ever smoker (which combines the UKB GWAS with the TAG GWAS; $n = 518,633$), and its P value in that GWAS is 3.92×10^{-18} .

3.4.5 GWAS of number of sexual partners

Our baseline GWAS protocol (described in **Supplementary Information section 2**) identified 118 lead SNPs associated with number of sexual partners. For reasons we explain below in this section, we exclude one of the 118 initially identified lead SNPs from our lead-SNP count for number of sexual partners, which leaves 117 number-of-sexual-partners lead SNPs. The strongest associations are located within, or in close proximity to, the genes *C14orf177* ($P = 4.61 \times 10^{-19}$) and *FURIN* ($P = 5.76 \times 10^{-17}$). Two lead SNPs are within, or in close proximity to, *CADM2* (rs2163971, $P = 4.6 \times 10^{-14}$; rs9856718, $P = 2.68 \times 10^{-8}$). 28 of the 117 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for general risk tolerance. 63 of the 117 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for at least one of the other main phenotypes we analyze, excluding the first PC of the risky behaviors.

Following the procedure described in **Supplementary Information section 2.10**, we determined that none of the lead SNPs we found (and the SNPs in LD with the lead SNPs, $r^2 > 0.1$) to be associated with number of sexual partners have been previously associated with any similar phenotype, and we therefore consider all of our associations with number of sexual partners to be novel (**Supplementary Table 3.2**).

Our baseline GWAS protocol identified 98 loci for number of sexual partners. However, for reasons we explain below in this section, we excluded one of these loci from the locus count, thus leaving 97 number-of-sexual-partners loci. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 158,435 SNPs that passed all GWAS quality control filters and that are located within the 98 loci. There were 88 genome-wide significant conditional associations, of which 82 are among the 118 lead SNPs. The six new conditional associations (COJO $P = 1.67 \times 10^{-12}$ – 2.55×10^{-8}), are all genome-wide significant in the GWAS (2.84×10^{-12} – 2.94×10^{-8}), but they are in the clumps of other lead SNPs. **Supplementary Table 3.2** lists the loci and reports the results of the conditional analysis.

We now discuss long-range LD and candidate inversion regions that contain number-of-sexual-partners lead SNPs. We emphasize again, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations. Five lead SNPs are located within the two long-range LD regions on chromosome 3 (~83.4 to 86.9 Mb) and chromosome 6 (~25.3 to 33.4 Mb) that are shared across general risk tolerance and all or almost all other GWAS (see **Supplementary Information section 3.2**). 25 lead SNPs are located within 14 candidate inversions. Furthermore, 23 lead SNPs are located within, or in strong LD with another variant within, 33 different 1000G structural variants (some lead SNPs are in strong LD with SNPs in multiple structural variants). All candidate inversions that contain lead SNPs for number of sexual partners, as well as these 1000G structural variants, are reported in **Supplementary Table 3.2**.

During the revision stage of this manuscript, some comments by a Referee prompted us to perform closer inspection of one of the 118 number-of-sexual-partners lead SNPs our baseline GWAS protocol had identified. The lead SNP in question, rs138394556 in locus no. 43 on chromosome 6

(at bp 30,652,782), is not in LD ($r^2 > 0.1$) with other SNPs in the main reference panel, and its very low MAF in our main reference panel (~ 0.0005) differs substantially from its MAF in the UKB (~ 0.148). rs138394556 was not directly sequenced (or did not pass QC filters) in the British-ancestry individuals in the 1000 Genomes phase 1 version 3 reference panel, and its MAF in the British-ancestry individuals in the 1000 Genomes phase 3 version 5^{90,91} ($n \sim 91$) is 0, similar to that in our main reference panel, and thus also very different from the MAF in the UKB.

In response to the Referee's comments, we first investigated the overall association support of all 865 initially identified lead SNPs from our seven GWAS that have no LD partners at a threshold of $r^2 > 0.6$. Out of 22 such lead SNPs, 10 are not in LD with any SNPs at $r^2 > 0.5$, three are not in LD with any SNPs at $r^2 > 0.4$, two are not in LD with any SNPs at $r^2 > 0.3$, and only a single lead SNP is not in LD with any SNPs at $r^2 > 0.1$ (rs138394556). Hence, for all the lead SNPs except rs138394556, there is LD support at lower r^2 thresholds.

There are two main possible causes of the observed MAF difference between our main reference panel and the UKB for rs138394556: (1) a genotyping or imputation error in the UKB data, or (2) a difference in MAF between British and European populations, combined with either a genotyping error in the 1000 Genomes data or non-representative sampling of the 1000 Genomes British individuals in a way that failed to capture the true MAF of rs138394556 in the entire British population.

To assess (1), we first investigated rs138394556 closely in the UKB. Although our GWAS analysis uses the imputed data, the SNP is directly genotyped with call rate of ~ 0.99 , and it passed all the internal QC filters of the UKB, described elsewhere³⁸. The genotyped MAF is ~ 0.14 , which matches the imputed data. Next, we investigated whether rs138394556's MAF varies across the UKB genotyping arrays and batches, as well as in the updated imputed data (referred to as version 3), released in March, 2018. Again, the MAF is highly similar across the arrays and batches, as well as in the updated imputed data (ranging from 0.128 to 0.151). These results, combined with the fact that our association analysis controls for batches and arrays, lead us to conclude that the association is not driven by array or batch differences. Furthermore, in the phenome-wide UKB analyses performed by the Neale lab (<http://www.nealelab.is/uk-biobank/>, accessed August 15, 2018), rs138394556 is suggestively associated with lifetime number of sexual partners in a smaller set of individuals ($P = 7.92 \times 10^{-6}$; $n = 296,609$); rs138394556 passed the Neale lab's QC filters (although they acknowledge their checks were rather rudimentary). Overall, we found no direct evidence of a genotyping or imputation error, and it is reassuring that the Neale lab also estimated an elevated level of association with number of sexual partners.

To investigate (2), we first checked rs138394556's MAF in the two study cohorts that provided the vast majority of the samples of British ancestry in our main reference panel, namely the UK IBD Genetics Consortium and the UK10K. We found that rs138394556 had not been directly sequenced in either cohort. We then reached out and heard back from investigators who work with cohorts with individuals of British ancestry (including ALSPAC, ELSA, the Fenland study, and UKHLS) to inquire whether rs138394556 had been genotyped in these cohorts and, if so, what its MAF was. The Fenland study ($n \sim 8,500$) is the only one of these cohorts in which rs138394556 has been directly genotyped. In the Fenland study, the genotyped MAF is ~ 0.13 and is thus very similar to that of the UKB. However, the Fenland study genotyped their participants with the UK Biobank Axiom genotyping array, the main array that was employed in the UKB. Thus, we cannot exclude the possibility that the genotyping arrays employed by the UKB fail to correctly genotype rs138394556.

Because of the strong overall association in the vicinity of rs138394556, which is clearly visible in the LocusZoom plot of locus no. 43, we have no reason to believe that the overall association in the region implicated by the SNP is the result of genotyping or imputation error. Therefore, we have decided to keep rs138394556 in the overall reported findings (e.g., in the Manhattan plot for number of sexual partners and in **Table S3.2**). However, because we cannot exclude the possibility that rs138394556 has been incorrectly genotyped, we conservatively decided to exclude rs138394556 from our count of the number-of-sexual-partners lead SNPs and loci. We added notes to highlight that rs138394556 was excluded from the lead-SNP counts when appropriate, to avoid confusion.

3.4.6 *GWAS of the first PC of the four risky behaviors*

We identified 106 lead SNPs associated with the first PC of the four risky behaviors. The strongest association is located in the gene *MAPT* on chromosome 17 (rs62062288, $P = 1.02 \times 10^{-29}$), and that gene contains lead SNPs for the GWAS of automobile speeding propensity, drinks per week, and number of sexual partners (but not for general risk tolerance, adventurousness, or ever smoker). The second strongest association is in *CADM2* (rs6790699, $P = 5.99 \times 10^{-27}$), our top gene associated with general risk tolerance, which also contains lead SNPs for all our main GWAS. The three next strongest associations are in *NCAM1* (rs2155290, $P = 4.11 \times 10^{-24}$, top locus for ever smoker), *ADH1B* (rs1229984, $P = 3.70 \times 10^{-22}$, top locus for drinks per week), and *FOXP1* (rs4676964, $P = 7.80 \times 10^{-18}$). This latter gene (*FOXP1*) is also strongly associated with the phenotypes automobile speeding propensity and number of sexual partners.

18 of the 106 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for general risk tolerance. 89 of the 106 lead SNPs are also lead SNPs, or in LD with a lead SNP ($r^2 > 0.1$), for at least one of the other main phenotypes we analyze. We note that a high level of overlap with the results of the GWAS of the four risky behaviors was to be expected, given that the first PC of the risky behaviors was obtained from a principal component analysis of the four risky behaviors.

We know of no previous GWAS of the first PC of the risky behaviors (or of similar phenotypes), and we consider all of our lead SNPs to be novel associations (**Supplementary Table 3.2**).

We identified 89 loci (**Supplementary Information section 2.8**) for the first PC of the risky behaviors. We also performed a conditional analysis with GCTA (COJO)⁵⁹ (**Supplementary Information section 2.8**) with the 120,227 SNPs that passed all GWAS quality control filters and that are located within the 89 loci. There were 84 genome-wide significant conditional associations, of which 79 are among the 106 lead SNPs. The five new conditional associations (COJO $P = 6.42 \times 10^{-12}$ – 3.76×10^{-9}), are all genome-wide significant in the GWAS (1.83×10^{-11} – 3.22×10^{-9}), but they are in the clumps of other lead SNPs. **Supplementary Table 3.2** lists the loci and reports the results of the conditional analysis.

We now discuss long-range LD and candidate inversion regions that contain first-PC lead SNPs. We emphasize again, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations. Seven of the 106 lead SNPs are located within the long-range LD region on chromosome 3 (~83.4 to 86.9 Mb), and three lead SNPs are located within the long-range LD region on chromosome 6 (~25.3 to 33.4 Mb), both shared with the general-risk-tolerance GWAS and all or most of our other main GWAS (see **Supplementary Information section 3.2**). One additional lead SNP is located within a long-range LD region on

chromosome 2 (~134.7 to 138.2 Mb). 20 lead SNPs are located within 14 candidate inversions. Of these, none is located within the chromosome 8 candidate inversion spanning ~7.89 to 11.8 Mb. The strongest association with the first PC of the risky behaviors within that candidate inversion is rs2898249 ($P = 1.27 \times 10^{-7}$), which we consider suggestive. Also, as can be seen from **Extended Data Fig. 3.4**, the general level of association within the candidate inversion on chromosome 8 (~7.89 to 11.8 Mb) is more similar to the phenotypes for which there are lead SNPs in the candidate inversion than to drinks per week (for which the candidate inversion does not contain any lead SNPs). Furthermore, 20 lead SNPs are located within, or in strong LD with another variant within, 24 different 1000G structural variants (some lead SNPs are in strong LD with SNPs in multiple structural variants).

3.4.7 Long-range LD regions and candidate inversions that contain lead SNPs for the supplementary GWAS phenotypes

Our seven main GWAS identified lead SNPs located in eight long-range LD regions and 44 candidate inversions (defined in **Supplementary Information section 2.9**). In **Supplementary Information section 3.2**, we focused on the two long-range LD regions and on the three candidate inversions that contain lead SNPs for general risk tolerance and for all or most of our six other GWAS phenotypes, and above in this subsection (**Supplementary Information section 3.4**) we have briefly presented the long-range LD regions and candidate inversions that contain lead SNPs for each of the six supplementary phenotypes. We now focus on three candidate inversions that are notable because they contain lead SNPs for four of the six other GWAS (but not for general risk tolerance). The remaining 37 identified candidate inversions and the remaining six long-range LD regions have little overlap across the GWAS and are reported in **Supplementary Tables 3.1** and **3.2**. We emphasize again, however, that the exact numbers of lead SNPs within the long-range LD and candidate inversion regions should be interpreted with caution because many of the lead SNPs in those regions are not conditional associations.

The first of the three additional candidate inversions that are notable because they are shared across four supplementary GWAS is located on chromosome 11 (~112.5 to 113.5 Mb). It contains one lead SNP for drinks per week, four for ever smoker, one for number of sexual partners, and one for the first PC of the risky behaviors. The strongest association with general risk tolerance within the candidate inversion is rs78168664 ($P = 1.47 \times 10^{-4}$). The GWAS Catalog database⁶⁵ reports a few previous phenotypes with associations in the region, which include bone ultrasound measurement, cardiac muscle measurement, and gut microbiota.

The second additional candidate inversion is on chromosome 16 (~12.0 to 14.8 Mb) and contains one lead SNP for drinks per week, one lead SNP for ever smoker, one lead SNP for number of sexual partners, and one lead SNP for the first PC of the four risky behaviors. The strongest association with general risk tolerance within the candidate inversion is rs2866323 ($P = 1.36 \times 10^{-5}$). The GWAS Catalog database⁶⁵ reports notable previous associations with traits such as age at menarche, human standing height, and schizophrenia.

The third candidate inversion is on chromosome 17 (~43.6 to 44.3 Mb) and contains one lead SNP for automobile speeding propensity, one for drinks per week, one for number of sexual partners, and two lead SNPs for the first PC of the risky behaviors. The strongest association with general risk tolerance within the candidate inversion is rs2866323 ($P = 1.86 \times 10^{-4}$). The candidate inversion covers only four genes, including the gene *MAPT* (~43.9 to 44.1 Mb on chromosome 17). SNPs

in and around *MAPT* have previously been associated with many neurodegenerative disorders, including Alzheimer's disease, Parkinson's disease and frontotemporal dementia⁷³. However, most of those previous associations are located outside the candidate inversion on chromosome 17 (~43.6 to 44.3 Mb), and *MAPT* is not significant after Bonferroni correction in the MAGMA gene analysis (**Supplementary Table 12.3**).

3.4.8 *GWAS Catalog lookup of the lead SNPs from the supplementary GWAS*

To investigate the overlap of our six supplementary GWAS phenotypes with previous GWAS of other phenotypes, we performed a lookup for each of the lead SNPs (and the SNPs in LD, $r^2 > 0.6$) in the NHGRI-EBI GWAS Catalog database⁶⁵, as described in **Supplementary Information section 2.11**. We report the results in **Supplementary Table 3.4**. For the six GWAS, we find in total 939 overlaps distributed across 130 lead SNPs. Since the six phenotypes are genetically correlated, they tend to share lead SNPs (or the lead SNPs for each phenotype are in LD, $r^2 > 0.1$), and as expected we find similar overlaps with other traits across the six phenotypes. Each of the six phenotypes have at least one overlap with all of the following phenotypes or phenotype categories: schizophrenia; educational attainment or cognitive performance (e.g. intelligence, information processing speed, cognitive function); BMI (body mass index) (or related anthropometric traits); circulating lipids (e.g. triglycerides, cholesterol, lipids); blood pressure or coronary artery disease (including arterial stiffness); and lung disease (e.g. interstitial lung disease, COPD, asthma).

Most of the six GWAS also have lead SNPs that overlap with the following phenotypes or phenotype categories: autoimmune diseases (e.g. Crohn's, rheumatoid arthritis, psoriasis, vitiligo); various cancers (e.g. breast cancer, lung cancer) or tumor formation; brain volume; bipolar disorder; age at menarche; Alzheimer's disease (driven by the *MAPT*-locus located in a candidate inversion on chromosome 17, ~43.6 to 44.3 Mb); height; Parkinson's disease; and attention hyperactivity deficit disorder.

The amount of overlap with schizophrenia seems particularly striking: in total, 19 distinct lead SNPs from our six additional GWAS overlap with loci associated with schizophrenia. We also observed many overlaps between general risk tolerance and schizophrenia (**Supplementary Information section 3.1**).

As discussed in **Supplementary Information section 3.3**, we note that both the existence and the absence of overlaps between any two phenotypes should be interpreted with care, and that the existence of overlaps is not necessarily evidence of shared genetic architecture⁸⁷.

3.5 Sensitivity analyses of the BiLEVE and Axiom samples

During the revision stage, a Referee raised the important point that the GWAS results could be sensitive to the sampling scheme of the UK BiLEVE study, whose participants were selected on the basis of their lung function and smoking behavior and were genotyped with a different genotyping array than the other UKB participants (**Supplementary Information section 2.5**). As we explain in **Supplementary Information section 2.5**, we believe it is preferable to analyze the complete UKB as a single cohort. Nonetheless, to gauge the sensitivity of our results to that decision, we repeated our discovery GWAS of general risk tolerance and our GWAS of ever

smoker, following the same protocol as for our main GWAS (**Supplementary Information section 2**) but treating the UK BiLEVE and the UKB Axiom cohorts as two separate cohorts^y.

We compared the results of the two new GWAS with the corresponding baseline GWAS along two dimensions. First, with bivariate LD Score regression²⁴, we estimated the genetic correlations across the new and the baseline GWAS. For both general risk tolerance and ever smoker, we estimated genetic correlations of exactly unity ($\hat{r}_g = 1$, $SE = 7.381 \times 10^{-6}$ for general risk tolerance; $\hat{r}_g = 1$, $SE = 5.643 \times 10^{-7}$ for ever smoker), thus indicating that the results are virtually the same at the aggregate level across the genome.

Second, we compared the lead SNPs across the new and the baseline GWAS. For general risk tolerance, the new discovery GWAS identified 120 genome-wide significant lead SNPs. Of the 124 lead SNPs of the baseline discovery GWAS, 113 are exactly the same; seven change to a SNP in the same locus and in high LD ($r^2 = 0.95-0.99$); and four are no longer lead SNPs in the new discovery GWAS, with their P values increasing from $P = 3.90-4.86 \times 10^{-8}$ to $P = 5.11-5.79 \times 10^{-8}$, just above the genome-wide significance threshold. The new discovery GWAS did not result in any new lead SNPs that were not genome-wide significant in the baseline discovery GWAS.

For ever smoker, the new GWAS resulted in 214 genome-wide significant lead SNPs. Of the 223 lead SNPs of the baseline GWAS, 190 are exactly the same; 16 change to a SNP in the same locus and in high LD ($r^2 = 0.29-0.99$); two are instead tagged by a single lead SNP in the new GWAS ($r^2 = 0.13$ and 0.70); and 15 are no longer lead SNPs in the new GWAS, with their P values increasing from $P = 3.49-4.99 \times 10^{-8}$ to $P = 5.65-1.16 \times 10^{-7}$, just above the genome-wide significance threshold. In addition, there are seven new genome-wide significant lead SNPs, all of which were just above genome-wide significance threshold in the baseline GWAS (baseline $P = 5.15-8.42 \times 10^{-8}$).

In summary, for both general risk tolerance and ever smoker the results are not particularly sensitive to the decision of treating the UKB as a single cohort, rather than treating the UK BiLEVE and the UKB Axiom cohorts as two separate cohorts.

3.6 Multiple regression with individual-level genotype dosages

In response to a comment by a Referee, we conducted a multiple regression analysis with individual-level genotype-dosage data from the UKB, to verify that the results of the conditional analysis we conducted with GCTA (COJO) are robust. In that analysis, for each phenotype (except adventurousness, for which there is no UKB data), for each chromosome we regressed the phenotype on all the phenotype's lead SNPs located on the chromosome (all jointly included in the same regression) and the control variables from our baseline association analyses (i.e., the top 20 principal components of the genetic relatedness matrix, sex-specific birth-year fixed effects, and fixed effects for genotyping batches; **Supplementary Information section 2.3** and **Supplementary Table 2.2**). Because BOLT-LMM does not offer the option to jointly fit multiple SNPs (and because we are unaware of other linear mixed models software that account for the relatedness of individuals while still being computationally efficient in a large sample like the UKB), we estimated linear regressions with SNPtest v2.5.4beta3⁹²; and because we estimated linear regressions instead of linear mixed models, we excluded one individual from each pair of

^y Because of participant withdrawal, the new GWAS contain seven fewer participants than the initial GWAS; this should have a negligible effect on this sensitivity analysis.

individuals whose genetic relatedness exceeds 0.044. We report the results in **Supplementary Table 3.1** and **Supplementary Table 3.2**.

For general risk tolerance, 90 lead SNPs were significant in the COJO analysis we conducted using the summary statistics of our discovery GWAS ($n = 939,908$) and our main reference panel ($n = 17,774$; **Supplementary Information section 2.4.1**). Despite the considerably smaller sample of unrelated individuals in the UKB which we used for the multiple regression analysis ($n = 355,727$), 89 of the 90 SNPs have consistent signs and are significant at the 5% level in the multiple regression analysis. The remaining SNP (rs17573719) also has a consistent sign and is marginally non-significant, with a P value of 0.064.

For automobile speeding propensity, 30 lead SNPs were significant in the COJO analysis we conducted using the summary statistics of our GWAS ($n = 404,291$). Despite the smaller sample ($n = 334,176$) used in the multiple regression analysis, all 30 SNPs have consistent signs and are significant at the 5% level in the multiple regression analysis; 28 are significant at $P < 1 \times 10^{-4}$, and 15 at $P < 5 \times 10^{-8}$.

For drinks per week, 56 lead SNPs were significant in the COJO analysis we conducted using the summary statistics of our GWAS ($n = 414,343$). Despite the smaller sample ($n = 342,239$) used in the multiple regression analysis, all 56 SNPs have consistent signs and are significant at the 5% level in the multiple regression analysis; all 56 are significant at $P < 1 \times 10^{-4}$ and 28 are significant at $P < 5 \times 10^{-8}$.

For ever smoker, 162 lead SNPs were significant in the COJO analysis we conducted using the summary statistics of our GWAS ($n = 518,633$). Despite the considerably smaller sample ($n = 366,921$) used in the multiple regression analysis, all 162 SNPs have consistent signs and are significant at the 5% level in the multiple regression analysis.

For number of sexual partners, 82 lead SNPs were significant in the COJO analysis we conducted using the summary statistics of our GWAS ($n = 370,711$). Despite the smaller sample ($n = 306,161$) used in the multiple regression analysis, all 82 SNPs have consistent signs and are significant at the 5% level in the multiple regression analysis; 77 are significant at $P < 1 \times 10^{-4}$, and 41 at $P < 5 \times 10^{-8}$.

For the first PC of the four risky behaviors, 79 lead SNPs were significant in the COJO analysis we conducted using the summary statistics of our GWAS ($n = 315,894$). Despite the smaller sample ($n = 261,485$) used in the multiple regression analysis, all 79 SNPs have consistent signs and are significant at the 5% level in the multiple regression analysis; 77 are significant at $P < 1 \times 10^{-4}$, and 36 at $P < 5 \times 10^{-8}$.

Overall, the results of our COJO analysis are consistent with those of our multiple regression analysis with individual-level genotype-dosage data in the UKB.

4 Testing for population stratification

Population stratification can be an important source of bias in GWAS. As per our analysis plan³⁶, every cohort in our GWAS included the ten (or more) top principal components (PCs) of the genetic-relatedness matrix (GRM) in their analyses, and some also used mixed-linear models. These procedures should help control for population stratification, but some population stratification and confounding bias could still remain after these controls⁹³.

In this section, we report the results of three tests of population stratification that are based on different sets of assumptions. The first test is the LD Score intercept test⁵³, which allows us to quantify the amount of stratification that is present in our estimates. The second is a sign test that compares the signs of GWAS estimates to the signs of the estimates of a within-family (WF) GWAS in an independent replication cohort. Third, we conduct a regression test that compares both the signs and the magnitudes of the GWAS estimates to those of the WF GWAS.

We applied the first test to the summary statistics of the discovery and replication GWAS of self-reported general risk tolerance and to the summary statistics of the supplementary GWAS; we applied the second and third tests to the summary statistics of the discovery GWAS of self-reported general risk tolerance.

4.1 LD Score intercept test

The LD Score intercept test uses GWAS summary statistics for all measured SNPs. Unlike the Genomic Control (GC) method, which assumes that confounding bias (e.g., due to population stratification and cryptic relatedness) is responsible for inflation in the GWAS χ^2 statistics, the LD Score regression method can disentangle inflation that is due to true polygenic signal throughout the genome (which affects the slope of the LD Score regression) from inflation that is due to confounding biases such as cryptic relatedness and population stratification (which affects the intercept of the regression).

We used the LDSC software⁵³ to estimate the intercepts in LD Score regressions with the summary statistics of our discovery and replication GWAS of general risk tolerance. We also estimated LD Score regressions with the summary statistics from the supplementary GWAS.

For each phenotype, we used the “eur_w_ld_chr/” files of LD scores computed by Finucane *et al.*⁹⁴ and made available on <https://github.com/bulik/ldsc/wiki/Genetic-Correlation>, accessed on March 14, 2016. These LD scores were computed with genotypes from the European-ancestry samples in the 1000 Genomes Project; only HapMap3 SNPs with MAF > 0.01 were included in the LD Score regressions. Because GC will tend to bias the intercept of the LD Score regression downward, we did not apply GC to the summary statistics prior to estimating the LD Score regressions.

Extended Data Fig. 4.1 shows LD Score regression plots for our discovery and replication GWAS and for the supplementary GWAS.

For our discovery GWAS, we estimated a LD Score intercept of 1.040 ($SE = 0.012$); for the replication GWAS, we estimated a LD Score intercept of 1.002 ($SE = 0.069$). The mean χ^2 statistics for all the SNPs in the two LD Score regressions are 1.848 and 1.031, respectively. The mean χ^2 statistics reflect the average strength of the GWAS associations between the SNPs and each phenotype. Under the null hypothesis that there is no confounding bias and that the SNPs

have no causal effects on the phenotypes, the mean χ^2 statistics would be 1. Thus, mean χ^2 statistics greater than 1 indicate that some SNPs are associated with the phenotypes, either because they are in LD with causal SNPs or because of confounding bias.

One measure of stratification bias is given by the ratio $\frac{Intercept-1}{\chi^2-1}$, which describes the share of the inflation in the mean χ^2 statistic ($\overline{\chi^2}$) that is due to stratification. This ratio is 0.048 ($SE = 0.014$) for the discovery GWAS and 0.071 ($SE = 0.021$) for the replication GWAS. These estimates imply that only a small part of the observed inflation in the mean χ^2 statistics from our discovery and replication GWAS is accounted for by confounding bias (due to population stratification, cryptic relatedness, or other confounds) rather than polygenic signal. This suggests that the bulk of the inflation in the χ^2 statistics from the discovery and replication GWAS is attributable to true polygenic signal throughout the genome, and that population stratification is unlikely to be a major concern for the analyses we present in this paper.

Supplementary Table 4.1 contains the full set of LD Score regression results for the discovery GWAS, the replication GWAS, the meta-analysis of the discovery and replication GWAS, and the supplementary GWAS. Although the LD Score intercepts from these regressions are often significantly larger than 1, the share of inflation in $\overline{\chi^2}$ that is due to stratification remains small (ranging from 0.047 to 0.071), which allows us to conclude that confounding bias is likely to account for no more than a small part of the inflation in these GWAS's mean χ^2 statistics.

As mentioned in **Supplementary Information section 2.7**, rather than applying the usual GC correction, we followed the increasingly common practice of adjusting the standard errors of our GWAS estimates for the possible effects of population stratification by inflating them by the square root of our estimates of the intercepts from the LD Score regressions⁵³. For the discovery and the replication GWAS of general risk tolerance, the meta-analysis of the discovery and replication GWAS of general risk tolerance, and the GWAS of ever smoker, each of which combines several cohorts, we inflated the standard errors at the meta-analysis level only.

4.2 GWAS/WF GWAS sign test for the general-risk-tolerance GWAS

As a simple test of whether the results of our general-risk-tolerance GWAS are driven entirely by stratification or whether they capture some genetic signal, we performed a sign test that compares the signs of the estimates from our discovery GWAS of general risk tolerance (excluding all full siblings from the UKB cohort, as described below) to the signs of the estimates from within-family (WF) GWAS of general risk tolerance in the independent STR1, STR2, and UKB-siblings cohorts. The UKB-siblings cohort was defined in the same way as the full UKB cohort, but only includes individuals with at least one full sibling in the UKB. To avoid overfitting (i.e., to ensure that the two sets of signs originate from GWAS that were conducted using independent cohorts)⁹⁵, we reran our discovery GWAS after excluding all individuals with at least one full sibling in the UKB.

If the discovery GWAS estimates are driven by stratification, then they should be independent of the signs of the WF GWAS estimates (which are immune to stratification) and therefore the two sets of signs should only have a concordance of roughly 50%. A significantly higher degree of sign concordance would suggest that at least some of the signal from the GWAS comes from true genetic effects.

4.2.1 Background

We followed the method outlined in Okbay *et al.* (2016)¹⁸. Here, we summarize the main assumptions and procedures that are described in more detail in that paper.

We first let $\hat{\beta}_j$ denote the estimate corresponding to SNP j from the discovery GWAS excluding the full siblings from the UKB, and let $\hat{\beta}_{WF,j}$ denote the WF GWAS estimate corresponding to SNP j . The WF GWAS estimate is obtained in a sample of sibling pairs, from a regression of sibling differences in general risk tolerance on sibling differences in genotypes and in controls. Since WF regressions are not biased due to stratification, and under the assumption that the WF effect size of each SNP is equal to the true population effect size, we can decompose the GWAS and WF GWAS estimates as:

$$\begin{aligned}\hat{\beta}_j &= \beta_j + s_j + U_j \\ \hat{\beta}_{WF,j} &= \beta_{WF,j} + V_j,\end{aligned}$$

where β_j and $\beta_{WF,j}$ are the true underlying GWAS and WF GWAS parameters for SNP j , s_j is the bias due to stratification (defined to be orthogonal to β_j and U_j), and U_j and V_j are the error terms in the estimates due to sampling variation, with $E(U_j) = E(V_j) = 0$. Note that if $\hat{\beta}_j$ and $\hat{\beta}_{WF,j}$ are estimated in independent samples, then U_j and V_j will be independent.

Under the null hypothesis that the GWAS contains no genetic signal (i.e., $\beta_j = \beta_{WF,j} = 0$ for all j) the sign of $\hat{\beta}_{WF,j}$ will be random with equal odds of being positive or negative and will be independent of the sign of $\hat{\beta}_j$. This means that among a set of M independent SNPs, the number of SNPs that have concordant signs, denoted C , follows a binomial distribution:

$$C \sim \text{Binomial}(M, 0.5).$$

We can thus measure the observed sign concordance and use this known distribution to formally test the null hypothesis. We tested this against the one-sided alternative hypothesis that $\beta_{WF,j}$ and β_j are not equal to zero and that their estimates thus have concordant signs. We conducted one-sided tests, because there is no reason to suspect that the signs would be discordant.

4.2.2 The GWAS and WF GWAS data

For this analysis, WF GWAS estimates were obtained in the STR1 (STR-Twingene), STR2 (STR-SALTY), and UKB-siblings cohorts. These estimates were then combined using a sample-size weighted meta-analysis. The STR1, STR2, and UKB-siblings cohorts include 674, 680, and 16,330 sibling pairs with general-risk-tolerance data, respectively. This gave us a total WF sample size of 17,684 sibling pairs (35,368 individuals).

The GWAS estimates are those from the discovery GWAS excluding all full siblings from the UKB, with a total sample size of 901,908.

Not every SNP from the GWAS results was available in the WF samples. To maximize power, we restricted our SNPs for each sign test to those that were available in both the GWAS results and in the three WF cohorts. Additionally, to ensure the quality of the WF GWAS estimates, we restricted our SNPs to those with $\text{MAF} \geq 0.05$ in every WF cohort and with imputation quality (INFO) above

95% in every WF cohort and above 99% in the discovery GWAS. Applying these filters left 2,005,496 SNPs in the intersection of the discovery and WF GWAS.

Then, we used PLINK⁴³ to apply a clumping algorithm to the GWAS results with the 1000 Genomes phase 3 EUR reference panel^z to obtain a subset containing approximately independent SNPs selected based on their GWAS P values. The clumping algorithm was similar to the main clumping algorithm we used to identify the GWAS's lead SNPs (**Supplementary Information section 2.8.1**) but used primary and secondary P value thresholds of $P = 1$ (instead of $P = 5 \times 10^{-8}$ and 1×10^{-4}). (The secondary P value threshold of $P = 1$ implies that at each iteration, all unclumped SNPs correlated ($r^2 > 0.1$) with the selected SNP are assigned to that SNP's clump, and the primary P value threshold of $P = 1$ implies that the clumping procedure keeps clumping until all SNPs have been clumped.) This procedure selected 48,979 approximately independent SNPs with results available for both the GWAS and the WF GWAS. Importantly, the procedure did not choose an arbitrary set of SNPs but instead selected the most significant SNP among the remaining (unclumped) SNPs at each iteration, such that each of the 48,979 SNPs is the most significant SNP in its clump.

4.2.3 Bayesian estimation of the posterior distribution of the SNPs' true effect sizes

Following Okbay *et al.* (2016)¹⁸, we conducted simulations to benchmark the results of our sign tests. To do so, we first obtained estimates of the distribution of the SNPs' true effect sizes (the β_j 's).

We define $\beta_{j,\text{std}}$ as the coefficient from the regression where both the phenotype and the genotype have been standardized to have mean zero and unit variance. We further assume that the effect sizes are drawn from a mixture distribution of a Gaussian and a point mass at zero:

$$\beta_{j,\text{std}} \sim \begin{cases} N(0, \tau^2) & \text{with probability } \pi \\ 0 & \text{otherwise,} \end{cases}$$

where τ^2 is the variance of non-null SNPs and π is the fraction of non-null SNPs in our data. This distributional assumption implies that the variance of effect sizes is inversely proportional to the variance of the unstandardized genotypes^{aa}. By the Central Limit Theorem, we note that the estimation error of the GWAS estimate of $\beta_{j,\text{std}}$ is approximately normally distributed. We use σ_j^2 to denote the variance of this error and note that our assumptions imply that $\sigma_j^2 \approx 1/n$. This means the distribution of $\hat{\beta}_{j,\text{std}}$ is:

$$\hat{\beta}_{j,\text{std}} \sim \begin{cases} N(0, \sigma_j^2 + \tau^2) & \text{with probability } \pi \\ N(0, \sigma_j^2) & \text{otherwise.} \end{cases}$$

Because we have a closed-form distribution, we can use the discovery GWAS summary statistics to estimate its parameters: the probability of a SNP being null π and the variance of the non-null effect sizes τ^2 (as mentioned above, $\sigma_j^2 \approx 1/n$).

^z This smaller reference panel (compared to main reference panel) suffices for this analysis because we only used SNPs with $\text{MAF} \geq 0.05$ in every WF cohort.

^{aa} To see this, note that $\beta_{j,\text{std}} = \beta_j \cdot \text{Var}(SNP_j)$ (where SNP_j is the unstandardized genotype at SNP j and can take values 0, 1, or 2). Because we assume that the distribution of $\beta_{j,\text{std}}$ is the same for all SNPs, it follows that $\text{Var}(\beta_j)$ will be larger for SNPs whose unstandardized genotypes have lower variance.

As shown in Okbay *et al.* (2016)¹⁸, given these parameters, the posterior probability that SNP j with estimated effect size $\hat{\beta}_{j,\text{std}}$ is non-null is:

$$p_{\hat{\beta},j} = \frac{\frac{\pi}{\sqrt{\sigma_j^2 + \tau^2}} \phi\left(\frac{\hat{\beta}_j}{\sqrt{\sigma_j^2 + \tau^2}}\right)}{\frac{1-\pi}{\sigma_j} \phi\left(\frac{\hat{\beta}_j}{\sigma_j}\right) + \frac{\pi}{\sqrt{\sigma_j^2 + \tau^2}} \phi\left(\frac{\hat{\beta}_j}{\sqrt{\sigma_j^2 + \tau^2}}\right)}.$$

And the posterior distribution of non-null SNPs is:

$$(\beta_j | \hat{\beta}_j, \beta_j \neq 0) \sim N\left(\frac{\tau^2}{\sigma_j^2 + \tau^2} \hat{\beta}_j, \frac{\sigma_j^2 \tau^2}{\sigma_j^2 + \tau^2}\right).$$

Following Okbay *et al.*, we estimated the parameters π , τ^2 , and $p_{\hat{\beta},j}$ using the summary statistics from the discovery GWAS (excluding the full siblings in the UKB cohort), restricting to the subset of 2,005,496 SNPs in the intersection of the discovery and WF GWAS (and with $\text{MAF} \geq 0.05$ in every WF cohort and imputation quality (INFO) above 95% in every WF cohort and above 99% in the discovery GWAS). Our estimates of π and τ^2 were 0.52 and 2.38×10^{-6} , respectively, with $p_{\hat{\beta},j}$ computed at the SNP level from the above equation and ranging from 0.38 to 1 (with a mean of 0.57 and standard deviation of 0.18)^{bb}. This generated a posterior distribution of the true effect size of each SNP given its estimated effect size.

4.2.4 Simulations

We used a simulation procedure whereby we drew, for each of the M approximately independent SNPs, “true” effect sizes from the resulting posterior distributions as well as estimation errors, and we repeated the simulation 1,000 times.

We let $E(C)$ denote the expected number of concordant signs under the alternative hypothesis that the SNPs have effect sizes given by the true β_j ’s. The quantity $E(C)$ provides a benchmark for the sign test. If the actual fraction of concordant SNPs matches the expected fraction $E(C)/M$ and is significantly larger than 50%, then we can be reasonably confident that most of the GWAS estimates are not driven by stratification.

We estimated $E(C)$ using the following method. For each of the 1,000 simulations, we generated discovery and WF GWAS estimates for each SNP j by adding Gaussian noise to the “true” effect size β_j drawn from the posterior distribution. We obtained the following quantities:

$$\hat{\beta}_{GWAS,j} = \beta_j + \varepsilon_j \hat{\sigma}_{GWAS,j}$$

^{bb} We also estimated $p_{\hat{\beta},j}$ for our 124 lead SNPs, and obtained a range of 0.9996 to 1 (compared to the 0.998 to 1 range reported in **Supplementary Information section 5.2.1**). These ranges differ slightly because our estimates of π and τ^2 were obtained using different sets of SNPs. In both cases, all lead SNPs have posterior probabilities of being causal near 1.

$$\hat{\beta}_{WF,j} = \beta_j + \varepsilon_j \hat{\sigma}_{WF,j},$$

where ε_j and δ_j are independent draws from a standard normal distribution and $\hat{\sigma}_{GWAS,j}$ and $\hat{\sigma}_{WF,j}$ are the standard errors of the coefficients for SNP j from the discovery and WF GWAS, respectively.

Let \hat{C}_k be the number of SNPs with matching discovery and WF GWAS signs in simulation k . We obtained our estimate of $E(C)$ by averaging \hat{C}_k across the 1,000 simulations:

$$\hat{E}(C) = \frac{1}{1000} \sum_{k=1}^{1000} \hat{C}_k.$$

In addition, we estimated the standard deviation of C by using the formula for the sample standard deviation^{cc}:

$$\widehat{SD}(C) = \sqrt{\frac{1}{999} \sum_{k=1}^{1000} (\hat{C}_k - \hat{E}(C))^2}.$$

4.2.5 Results of the sign test

Supplementary Table 4.2 reports the results of a range of sign tests we conducted. We conducted four sign tests, all of which compared the signs of the estimates from the discovery GWAS of risk tolerance (excluding all full siblings from the UKB) to the signs of the estimates from the meta-analysis of the WF GWAS in the UKB-siblings, STR1, and STR2 cohorts. Before each sign test, we pruned the list of independent clumped SNPs based on the P values obtained in the discovery GWAS. Each sign test corresponds to one of the following four P value cutoffs: 1 (all SNPs included), 0.05, 0.005, and 0.0005.

As can be seen from **Supplementary Table 4.2**, we strongly reject the null hypothesis of no sign concordance for all of the sign tests. All four sign tests are significant at the 5×10^{-10} level.

Note that all of the sign tests we conducted are well powered to reject the null hypothesis of no sign concordance. Based on these sign tests, we can be reasonably confident that at least some of the GWAS estimates are not driven by population stratification.

Supplementary Table 4.2 also reports the expected number of concordant signs for each of the four sign tests that were conducted. For each test, the observed number of concordant signs is similar to, but nonetheless significantly smaller than, the expected number of concordant signs predicted by the simulation.

This discrepancy between the observed and expected number of concordant signs could be attributable to several factors. First, of course, there could be population stratification (although the LD Score regression results suggest that there is unlikely to be much stratification); because

^{cc} This procedure should yield unbiased estimates of the expected value and the standard deviation of C , conditional on the estimated posterior distribution of the true effect sizes. Because we used an independent set of SNPs, both the expected value and the standard deviation of C are additive functions of the SNP-level probabilities of sign concordance, and this holds even though the simulation does not take the LD between the SNPs into account.

the simulation assumes no population stratification, the presence of population stratification would lead to fewer observed concordant signs than expected. Second, the discrepancy could be the result of our assumption, under the null hypothesis and in the simulation, that the SNPs' true effect sizes are the same in the discovery and WF GWAS (i.e., that $\beta_j = \beta_{WF,j}$). This assumption could fail for several reasons. For example, under positive assortative mating, the magnitude of the true GWAS effects would be larger than that of the true WF GWAS effects. The true discovery and WF GWAS effects would also be different if, for instance, parents were successful in partly counteracting differences in genetic propensity for risk tolerance and in making their children more similar to each other in terms of their risk tolerance; they would also be different if parental risk tolerance had sizeable effects on their children's risk tolerance via the rearing environment (as recently documented in the case of educational attainment⁹⁶).

4.3 Within-family regression test for the general-risk-tolerance GWAS

The above sign test compares the signs of the estimates of the discovery and WF GWAS of general risk tolerance. In this section, we introduce and conduct a third test, the within-family regression test, which allows us to compare both the signs and the estimates of the discovery and WF GWAS of general risk tolerance. This in turn allows us to draw some insights about some of the reasons why there is a discrepancy between the observed and the expected number of concordant signs in the sign test.

4.3.1 Background

Lee *et al.* (2017)⁹⁷ (in Supplementary Information section 2.8) provides an in-depth description of the within-family regression test. Following the notation in **Supplementary Information section 4.2.1**, we define the parameter

$$m_c \equiv \frac{\text{Cov}(\beta_j, \beta_{WF,j})}{\text{Var}(\beta_j) + \text{Var}(s_j)},$$

where β_j is the true effect size for SNP j in the discovery GWAS of general risk tolerance, $\beta_{WF,j}$ is the true effect size for SNP j in the WF GWAS of general risk tolerance, and s_j is the stratification bias for SNP j .

This m_c parameter can be interpreted in light of a few special cases. First, if the true discovery and WF GWAS effects are identical (i.e., if $\beta_j = \beta_{WF,j}$; this implies that the genetic correlation between the discovery and WF GWAS is unity and the effects are of equal magnitudes), then m_c reduces to $\frac{\text{Var}(\beta_j)}{\text{Var}(\beta_j) + \text{Var}(s_j)}$, which is the share of variance in the discovery GWAS estimates that is due to true signal. This quantity converges to one if there is no stratification and to zero if the discovery GWAS estimates are entirely driven by stratification.

The second special case occurs if there is no population stratification (i.e., if $\text{Var}(s_j) = 0$). Then, the parameter reduces to $\frac{\text{Cov}(\beta_j, \beta_{WF,j})}{\text{Var}(\beta_j)}$, which is the slope of the regression of the WF GWAS estimates on the discovery GWAS estimates.

4.3.2 Methods

We used the same consistent estimator of m_c as Lee *et al.* (2017)⁹⁷:

$$\widehat{m}_c \equiv \frac{\widehat{\text{Cov}}(\hat{\beta}_{GWAS,j}, \hat{\beta}_{WF,j})}{\widehat{\text{Var}}(\hat{\beta}_{GWAS,j}) - \widehat{\sigma}_{GWAS}^2}.$$

Our estimator for $\widehat{\sigma}_{GWAS}^2$ is $\widehat{\sigma}_{GWAS}^2 \equiv \frac{1}{J} \sum_{j=1}^J \widehat{\sigma}_{GWAS,j}^2$, where J is the number of SNPs in the full discovery GWAS.

To compute $\widehat{\text{Cov}}(\hat{\beta}_{GWAS,j}, \hat{\beta}_{WF,j})$ and $\widehat{\text{Var}}(\hat{\beta}_{GWAS,j})$, we applied a pruning procedure to select a set of approximately independent SNPs that are not prioritized by P value. To do this, we used PLINK's --indep-pairwise option with the following parameters: a window size of 50 SNPs, a window shift of 5 SNPs, and a pairwise r^2 threshold of 0.1. This yielded 51,473 approximately independent SNPs (unlike the 48,979 SNPs we obtained from the clumping procedure in **Supplementary Information section 4.2.2**, these 51,473 SNPs were not selected based on their P values.) To compute $\widehat{\text{Cov}}(\hat{\beta}_{GWAS,j}, \hat{\beta}_{WF,j})$, we used the same WF GWAS as in the sign test (**Supplementary Information section 4.2.2**).

We obtained the 95% confidence interval for \widehat{m}_c by bootstrapping the 51,473 approximately independent SNPs. We used the 25th and 975th smallest estimates of \widehat{m}_c out of 999 bootstrap draws as the lower and upper bounds of the confidence interval. We also calculated the bootstrap standard error by taking the standard deviation of the 999 bootstrap draws.

4.3.3 Results

We estimated $\widehat{\text{Cov}}(\hat{\beta}_{GWAS,j}, \hat{\beta}_{WF,j}) = 1.05 \times 10^{-6}$, $\widehat{\text{Var}}(\hat{\beta}_{GWAS,j}) = 2.16 \times 10^{-6}$, and $\widehat{\sigma}_{GWAS}^2 = 1.11 \times 10^{-6}$. Thus, our estimate of \widehat{m}_c was 1.00, with a bootstrap confidence interval of 0.87 to 1.11. (Following Lee *et al.* (2017), we also obtained standard errors using the block-jackknife procedure introduced by Bulik-Sullivan *et al.*⁵³. In our case, each block consists of a set of approximately 50 adjacent SNPs. We obtained a jackknife standard error of 0.10, which is larger than the bootstrapped standard errors of 0.06; thus, the bootstrap confidence interval may be slightly too tight, likely due to the fact that the 51,473 approximately independent SNPs are not fully independent of one another.)

In the first special case where $\beta_j = \beta_{WF,j}$, this estimate of m_c suggests, with roughly 95% confidence, that at most 13% of the variation in the GWAS estimates is due to stratification. This is consistent with the results from LD Score regression, which imply that about 5% of the variation in the GWAS estimates is due to stratification.

However, given the sign test results, it is likely that the assumption $\beta_j = \beta_{WF,j}$ does not hold exactly. The confidence interval for m_c is consistent with moderate differences in the magnitude of the GWAS estimates and/or with imperfect genetic correlation between the discovery and WF GWAS. If we assume, as implied by the results from LD Score regression, that 5% of the variance in the discovery GWAS estimates is due to stratification, then the 95% CI of 0.87 to 1.11 is consistent with an effect-size ratio ($\beta_{WF,j}/\beta_j$) of as low as 0.92 ($= 0.87/0.95$) and as high as 1.17 ($= 1.11/0.95$).

Small differences between β_j and $\beta_{WF,j}$ suggest that general risk tolerance is unlikely to be subject to much assortative mating; that parents are unlikely to actively enforce similarity in risk preferences among their children (or to be successful in doing so); and that parental risk tolerance is unlikely to have large effects on children's risk tolerance via the rearing environment (unlike what was recently documented in the case of educational attainment⁹⁶).

4.4 Discussion

We have presented the results of three tests of population stratification that rely on different sets of assumptions. The LD Score intercept test relies on stronger assumptions and allows us to quantify how much population stratification is present in our estimates. Our results from this test—for our discovery and replication GWAS of general risk tolerance and for our supplementary GWAS—imply that only a small part of the observed inflation in the mean χ^2 statistics is likely to be accounted for by confounding bias rather than polygenic signal.

Our results from the second test, the sign test, allow us to strongly reject the null hypothesis that the coefficients from our discovery GWAS of general risk tolerance are driven by stratification. All four sign tests are significant at the 5×10^{-10} level.

Lastly, the within-family regression test takes both the signs and the magnitudes of the GWAS estimates into account, and allows us to establish bounds on the amount of stratification and on the differences between the WF and discovery GWAS effects. From this test, we conclude that SNP effects are likely similar between and within families, and that population stratification is unlikely to be a major source of bias.

In sum, all three tests allow us to conclude that our results are unlikely to be driven by population stratification. We can be reasonably confident that the bulk of the variation in the GWAS estimates is attributable to true polygenic signal.

5 Replication of the general-risk-tolerance lead SNPs and maxFDR calculation

To assess the credibility of the results of our discovery GWAS of self-reported general risk tolerance, we attempted to replicate the associations of the lead SNPs from that GWAS in an independent replication GWAS of self-reported general risk tolerance.

As described in **Supplementary Information section 2.1**, the replication GWAS included 10 cohorts, with a combined sample size of 35,445. To assess the replicability of the lead SNPs from our discovery GWAS, we followed the procedure outlined in Supplementary Information section 1.8 of Okbay *et al.* (2016)¹⁶ and conducted binomial tests to assess whether the associations of the lead SNPs from our discovery GWAS replicate in an independent replication GWAS. We conducted two binomial tests. First, we conducted a binomial sign test to assess whether the directions (i.e., the signs) of the effects of the lead SNPs are concordant across the discovery and the replication GWAS. Second, we conducted a binomial test to assess whether a larger fraction of the lead SNPs are significant at the 5% level in one-sided tests (i.e., with concordant signs and significant at the 10% level on two-sided tests) in the replication GWAS, relative to what can be expected by chance.

We benchmarked these results against a plausible alternative hypothesis, where we predicted the number of concordant signs and significant SNPs in the replication GWAS given a posterior estimate of the true distribution of effect sizes. This allowed us to determine whether the results of the two aforementioned binomial tests match what we would expect if the lead SNPs were all true positives. Lastly, to ensure that our replication results are not biased due to overfitting, we assessed the extent of sample overlap across our discovery and replication GWAS, and tested the robustness of our results to excluding the UKHLS cohort from the replication GWAS.

In addition, to provide some reassurance about the reliability of the results of our seven main GWAS, we calculated the “maxFDR”, an upper bound on the false discovery rate (FDR), for each GWAS.

5.1 Methods

5.1.1 Constructing the set of lead and proxy-lead SNPs

We used the 124 independent lead SNPs from our discovery GWAS of general risk tolerance to construct a set of lead and proxy-lead SNPs that are available in the replication GWAS summary statistics. We first identified the subset of the 124 lead SNPs that were directly available in the replication GWAS summary statistics and had a sample size of at least one-half the maximum sample size in that GWAS. We identified 122 such SNPs. Next, for each of the remaining two SNPs, we determined whether there exists a suitable “proxy-lead SNP” that satisfies three conditions: (1) the SNP is in high LD ($r^2 > 0.8$) with the original SNP (we used PLINK⁴³ and our main reference panel to compute LD); (2) the SNP is available in the summary statistics of both the discovery GWAS and replication GWAS; and (3) the SNP has a sample size of at least one-half the maximum sample size in the replication GWAS. If more than one proxy-lead SNP satisfied these three condition for one original SNP, we selected the one in highest LD with the original SNP as the proxy-lead SNP for the analyses. We identified one such proxy-lead SNP. We

combined that proxy-lead SNP with those that were directly available in the replication GWAS to create the set of 123 lead and proxy-lead SNPs.

5.1.2 *Binomial replication tests*

We conducted two binomial tests of the null hypothesis that none of the lead SNPs are associated with risk tolerance. Rejecting this null provides evidence that our lead SNPs contain at least some truly associated SNPs. Under the null, we would expect that 50% of the lead SNPs have concordant signs and 5% are significant at the 5% level on the one-sided tests (i.e., significant at the 10% level, with concordant signs) in the replication GWAS. For both binomial tests, we used one-sided tests of the null hypothesis because we are specifically interested in testing for a larger share of concordant or significant SNPs relative to the null share.

Because the lead SNPs are approximately independent (pairwise $r^2 < 0.1$), the number of lead SNPs that have concordant signs or that are significant in the replication GWAS under the null can be modeled as a series of coin flips, where the probability of a “success” is 0.5 for sign concordance and 0.05 for the one-sided tests at the 5% level of significance. It follows that under the null hypothesis, k (which we define as the total number of concordant or significant SNPs) is distributed $k \sim \text{Binomial}(M, \pi)$, where M is the total number of lead SNPs and π is the probability of encountering a concordant or significant SNP (i.e., 0.5 for sign concordance or 0.05 for the one-sided tests at the 5% level of significance). Knowing the distribution of k allows us to perform tests of the null hypothesis. The results of these tests are presented below.

5.1.3 *Expected replication record*

In addition to the test of the null hypothesis described above, we also benchmarked our replication record against an estimate of a plausible replication record.

As for the sign test in **Supplementary Information section 4.2**, we followed the procedure outlined in Okbay *et al.* (2016)¹⁶ and conducted a Bayesian analysis to obtain estimates of the posterior distribution of the SNPs’ true effect sizes (the β_j ’s), given their GWAS estimates. That procedure is described in more detail in **Supplementary Information section 4.2** and in Okbay *et al.* (2016)¹⁶.

To compute the expected sign concordance and its variance under the posterior distribution of the SNPs’ true effect sizes, we proceeded as in **Supplementary Information section 4.2**, except that we used the set of lead and proxy-lead SNPs (instead of the ~50,000 approximately independent SNPs) and replaced the within-family GWAS by the replication GWAS. Our estimates of π and τ^2 were 0.29 and 2.02×10^{-6} , respectively (they differ from those reported in **Supplementary Information section 4.2** because they were estimated using all SNPs in the discovery GWAS and summary statistics that were inflated by the square root of the LD Score regression intercept).

To compute the expected replication record and its variance at the 5% level under the posterior distribution of the SNPs’ true effect sizes, we proceeded as in **Supplementary Information section 4.2**, except that we replaced \hat{C}_k with \hat{R}_k , where \hat{R}_k is defined as the number of SNPs in simulation k where the simulated replication effect is a significant replication of the simulated discovery effect.

5.2 Replication results

5.2.1 *Baseline replication results*

Out of the 123 lead or proxy-lead SNPs, 94 have concordant signs in the replication GWAS. Under the null hypothesis that the lead SNPs from our discovery GWAS are all null, we would expect 50% of the SNPs (i.e., 61.5 SNPs) to have concordant signs in the replication GWAS. Based on the results from our binomial test of sign concordance, we strongly reject the null hypothesis of 50% concordance ($P = 1.7 \times 10^{-9}$). Our expected replication record is that 95.9 ($SD = 4.6$) of the 123 SNPs would have matching signs, which matches our actual replication record very closely.

Out of the 123 lead or proxy-lead SNPs, 23 are significant at the 5% level on one-sided tests (i.e., significant at the 10%, with concordant signs) in the replication GWAS. Under the null hypothesis that the lead SNPs from our discovery GWAS are all null, we would expect 5% of the SNPs (i.e. 6.15 SNPs) to reach significance at the 5% level on the one-sided tests in the replication GWAS. We strongly reject this null hypothesis ($P = 4.5 \times 10^{-8}$). Consistent with this, our expected replication record is that 24.8 ($SD = 4.4$) SNPs would be significant at the 5% level, which again matches our actual replication record very closely.

The replication record of our lead and proxy-lead SNPs is shown in **Extended Data Fig. 5.1**, and the summary statistics for the 123 lead or proxy-lead SNPs are reported in **Supplementary Table 3.1**.

In sum, our actual replication record matches our expected replication record for both the sign and significance binomial tests. Moreover, both binomial tests strongly reject the null hypothesis that none of our lead SNPs are true associations. We also note that under our Bayesian model of true effect sizes, the posterior probability that a SNP is causal is above 99.8% for all of our 124 lead SNPs. Coupled with our empirical replication results, this suggests that most if not all of the lead SNPs we identified are true positives.

5.2.2 *Robustness to potential sample overlap between the discovery and replication GWAS*

Potential overlap between our discovery and replication samples is a concern for the validity of our replication results. Following a comment by a Referee, we conducted additional analyses to assess the extent of sample overlap across our discovery and replication GWAS and between the UKB and the UKHLS, and to assess the robustness of our replication results to excluding the UKHLS cohort from the replication GWAS.

First, to assess the extent of effective overlap between our discovery and replication GWAS samples, we estimated the intercept in a bivariate LD Score regression²⁴ using the summary statistics of our discovery and replication GWAS of general risk tolerance. According to theory, the intercept is equal to $\rho N_S / \sqrt{N_1 N_2}$, where ρ is the phenotypic correlation among the N_S individuals included in both samples (we assumed $\rho = 1$), and N_1 and N_2 are the sizes of the discovery and replication samples (so here $N_1 = 939,908$ and $N_2 = 35,445$). From this, we obtained an estimate of N_S : $\hat{N}_S = 602$ ($SE = 1,314$). This estimate is not significantly different from zero.

Second, following the Referee’s suggestion, we assessed the extent of sample overlap between the UKB and the UKHLS cohort. The UKB comprises a large fraction of our discovery sample, and the UKHLS cohort was included in our replication GWAS. Because both cohorts are comprised of individuals living in the UK, sample overlap could be a concern. Using bivariate LD Score regression, we obtained an estimate $\hat{N}_S = 615$ ($SE = 328$). The estimate is almost significant at the 5% level ($P = 0.07$) and is relatively large relative to the size of the UKHLS cohort ($n = 5,898$), thus suggesting there may be some effective sample overlap between the UKB and the UKHLS cohorts.

To verify that this effective sample overlap between the UKB and the UKHLS cohort does not bias our replication results, we reran the replication analyses after excluding the UKHLS cohort from our replication GWAS. We obtained a new expected sign concordance of 93.7 SNPs ($SD = 4.7$) and an expected 22.6 significant SNPs ($SD = 4.3$) at the 5% level. Our actual replication record matches this closely, with 91 SNPs with concordant signs and 21 SNPs replicating at the 5% level.

In summary, there appears to be no more than minimal sample overlap between our discovery and replication GWAS, and our replication results do not appear to be sensitive to overlap between the UKB and UKHLS.

5.3 maxFDR calculation

To provide additional reassurance about the reliability of our general-risk-tolerance lead SNPs, and some reassurance about the reliability of the lead SNPs from our six supplementary GWAS, we followed the methodology in Turley *et al.* (2018)⁷¹ and calculated the “maxFDR” for each GWAS. maxFDR is an upper bound on the false discovery rate (FDR) for the results of a GWAS. The key assumption of the maxFDR calculation is that the effect sizes for the phenotype of interest follow a spike-and-slab distribution. Under this assumption, the FDR is a function of the fraction of null SNPs (π_{null}), the variance of the effect sizes for non-null SNPs ($\sigma_{nonnull}^2$), and the sample size (N) (see Section 1.4.3 of Turley *et al.*’s Supplementary Information). For any given value of π_{null} , the value $\sigma_{nonnull}^2$ is determined by N and the mean χ^2 -statistic of the GWAS. Using the observed values of N and mean χ^2 -statistic across all the SNPs analyzed in the GWAS, we can therefore calculate the FDR for any value of π_{null} . The maxFDR is defined as the maximum theoretical FDR over a range of values for π_{null} . We searched over the range $\pi_{null} \in [0.02, 0.98]$ and evaluated the FDR at every point in the interval at 0.02 unit increments. To calculate maxFDR for the results of one of our seven GWAS, we used the MTAG software⁷¹ and passed into the software only our results for that GWAS (and not the results of other GWAS).

For general risk tolerance, we estimated that the maxFDR is 2.02×10^{-4} , which is achieved when $\pi_{null} = 0.26$. (Our maximum-likelihood estimate of π_{null} for risk tolerance, estimated with the procedure described in **Supplementary Information section 4.2.3** but using the summary statistics of all SNPs in the GWAS and substituting $1 - \pi_{null}$ for π (since $\pi_{null} = 1 - \pi$), is actually 0.71. Using this value for π_{null} yields a lower FDR of 4.10×10^{-5} .) The strong replication record of our risk tolerance lead SNPs (see **Supplementary Information section 5**) gives us additional confidence that the false positive rate among our risk tolerance results is likely to be very low.

For the six supplementary GWAS, given the very large sample sizes we used in our study, we anticipate that the rate of false positives is likely to be extremely low at the genome-wide significance threshold of 5×10^{-8} . Consistent with this, we calculated maxFDR’s of 1.23×10^{-4} , 1.22×10^{-3} , 7.08×10^{-4} , 1.50×10^{-4} , 3.00×10^{-4} , 3.06×10^{-4} for adventurousness, automobile speeding

propensity, drinks per week, ever smoker, number of sexual partners, and the first PC, respectively. These low maxFDR estimates give us additional confidence that the results of our six supplementary GWAS are robust.

6 Estimation of genome-wide SNP heritability

Risk tolerance (both self-reported and experimentally elicited) has been found to be moderately heritable in twin studies, with heritability estimates ranging from 20% to 60%^{2,7,8}. In this section, we employ three different methods to obtain estimates of the SNP heritability of our primary and supplementary GWAS phenotypes. A phenotype's SNP heritability is the fraction of the phenotype's variance that is accounted for by the additive genetic effects of a set of SNPs.

6.1 Methods to estimate genome-wide SNP heritability

We used three methods, GCTA⁹⁸, LD Score regression⁵³, and Heritability Estimator from Summary Statistics (HESS)⁹⁹, to estimate the genome-wide SNP heritability (h_G^2). The GCTA method estimates the heritability of a phenotype directly from the individuals' genotypic data, while the LD Score and HESS methods use GWAS summary statistics as inputs.

For comparability across phenotypes and methods, for the LD Score and HESS methods, we used summary statistics from the UKB GWAS only for all phenotypes except adventurousness. For the adventurousness phenotype, we only report estimates that were obtained using the LD Score regression and HESS methods using the 23andMe summary statistics, because this phenotype is not available in the UKB and we did not have access to the individual-level genotypic data from the 23andMe cohort (and so could not obtain estimates with the GCTA method).

We also computed HESS SNP heritability estimates using summary statistics from our seven main GWAS (and not only from GWAS conducted in the UKB or in the 23andMe cohort).

6.1.1 The GCTA method

The GCTA method is based on restricted maximum-likelihood estimation and uses the genetic relationship matrix (GRM) to estimate the SNP heritability. Under the assumptions discussed in Yang *et al.* (2011)⁹⁸, the method leads to unbiased estimates of the genome-wide SNP heritability. However, it is computationally intensive, and it is thus necessary to limit the number of SNPs and individuals included in the analysis in order to be computationally feasible. Therefore, we restricted the GCTA analysis to a random subset of 30,000 individuals out of the full sample from the discovery GWAS. We thereafter dropped one individual in each pair of individuals with a cryptic relatedness exceeding 0.025, to obtain a set of unrelated individuals. For comparability we used the same initial subset of 30,000 individuals for the GCTA estimation for all phenotypes, though the sample size varies slightly across phenotypes because of missing phenotypic observations. The final sample sizes for each phenotype are presented in **Supplementary Table 6.1**. In total 646,855 directly genotyped SNPs with MAF > 0.01 were included in the GCTA heritability estimation.

6.1.2 The LD Score regression method

Under the assumptions discussed in Bulik-Sullivan *et al.* (2015)⁵³, a SNP's GWAS χ^2 statistic is linearly related to its LD score, defined as the sum of the squared correlation coefficients between any single SNP and all the other SNPs. The slope of the LD Score regression (of the SNPs GWAS χ^2 statistics on their LD scores and an intercept) can be rescaled to obtain an estimate of the heritability explained by the SNPs included in the LD Score analysis by dividing the slope by the

sample size divided by the number of SNPs, i.e., by n/M . We used the “eur_w_ld_chr/” files of LD scores computed by Finucane *et al.*⁹⁴ and made available on <https://github.com/bulik/ldsc/wiki/Genetic-Correlation>, accessed on March 14, 2016. These LD scores were computed with genotypes from the European-ancestry samples in the 1000 Genomes Project. Only HapMap3 SNPs with MAF > 0.01 were included in the LD Score regression; for every phenotype, ~1.3 million SNPs were used for the LD Score heritability estimation. Since Genomic Control (GC) will tend to bias the intercept of the LD Score regression downward, we did not apply GC to the summary statistics prior to estimating the LD Score regressions.

6.1.3 The HESS method

The HESS estimator can be described in brief as an analytical variance decomposition method that, unlike the GCTA and LD Score regression methods, assumes that the SNP effect sizes are fixed effects rather than random effects. The method assumes that the SNPs are randomly distributed in the population and requires a pre-specified SNP covariance matrix as input. The SNP covariance matrix can be estimated in the sample of interest if individual genotypic data is available, or with an external reference panel such as the 1000 Genomes. As Shi *et al.*⁹⁹ show using simulations, heritability estimates from LD Score regression are sensitive to the true proportion of causal SNPs, and the HESS estimator yields more accurate heritability estimates than LD Score regression under a wider range of proportions of truly causal SNPs. We used the reference panel distributed with the HESS software for the calculation of the covariance matrix. That panel is the European subsample of the 1000 Genomes phase 3 version 5 reference panel, restricted to common variants (MAF > 0.05), which is the same as the reference panel used for the construction of the LD Scores^{dd}. For every phenotype, a total of ~4.9 million SNPs were used in the HESS heritability estimation. As with the LD Score regressions, we did not apply GC prior to estimating heritability with HESS.

6.2 Results

6.2.1 Results of genome-wide SNP heritability estimation

The results of the genome-wide SNP heritability estimations are reported in **Supplementary Table 6.1** and displayed in **Extended Data Fig. 6.1**. The estimated heritabilities of our primary and supplementary GWAS phenotypes range from 0.055 to 0.173.

For self-reported general risk tolerance, we obtained a GCTA heritability estimate of $\hat{h}_{GCTA}^2 = 0.085$ ($SE = 0.018$), a LD Score heritability estimate of $\hat{h}_{LD\ Score}^2 = 0.055$ ($SE = 0.002$), and a HESS heritability estimate of $\hat{h}_{HESS}^2 = 0.063$ ($SE = 0.003$). Of all estimated phenotype, the highest estimated heritability was for the first PC of the risky behaviors: $\hat{h}_{GCTA}^2 = 0.173$ ($SE = 0.025$), $\hat{h}_{LD\ Score}^2 = 0.114$ ($SE = 0.004$), and $\hat{h}_{HESS}^2 = 0.156$ ($SE = 0.004$).

The methods yield broadly consistent results. For all phenotypes, the heritability estimates are similar across the three methods, although the GCTA heritability estimates are generally the highest and the LD Score regression estimates are generally the lowest.

^{dd} While the same reference panel was used for the construction of the LD scores, as indicated above HapMap3 SNPs with MAF > 0.01 were included in the LD score regression.

We emphasize that, for general risk tolerance and ever smoker, our main GWAS also analyzed data from cohorts other than the UKB, so the heritability estimates we report in **Supplementary Table 6.1** and display in **Extended Data Fig. 6.1** are different from those that would have been obtained using the summary statistics from our main GWAS. **Table 1** reports HESS SNP heritability estimates obtained by using summary statistics from our main GWAS of the seven phenotypes.

6.2.2 *Relationship between genome-wide SNP heritability and number of lead SNPs across the seven main GWAS*

Following a suggestion by a Referee, we conducted an exploratory analysis of the relationship between the SNP heritability of our seven main phenotypes and the number of lead SNPs identified in the GWAS of each phenotype. We regressed the number of lead SNPs identified in each GWAS on the phenotype's SNP heritability, while controlling for the square root of the GWAS sample size. We note that because this regression includes only seven observations and includes a constant and two covariates, there are only four degrees of freedom, so statistical power is very limited. We used the HESS SNP heritability estimates from **Table 1** because these were estimated using the summary statistics from the full GWAS of each phenotype. We only included lead SNPs with a minor allele frequency (MAF) above 0.05, because the HESS heritability estimates were produced using only SNPs with MAF above 0.05. We estimated that a one-percentage-point increase in the SNP heritability is associated with 16.4 ($SE = 9.3$) additional discovered SNPs. However, this estimate is imprecise and not statistically significant ($P = 0.15$).

7 Genetic correlations

To assess whether the genetic variants that influence general risk tolerance tend to be the same and to have similar effect sizes as those that influence plausibly related phenotypes, we used bivariate LD Score regression²⁴ to estimate pairwise genetic correlations for autosomal SNPs between self-reported general risk tolerance and a number of pre-selected phenotypes. We also estimated the genetic correlation for general risk tolerance between males and females, between the discovery and replication GWAS, and between the 23andMe and UKB cohorts, and we compared the genetic and phenotypic correlations between general risk tolerance and our four main risky behaviors.

7.1 Methodology

Under some assumptions, bivariate LD Score regression²⁴ produces unbiased estimates of genetic correlation, even in the presence of sample overlap. Under these assumptions the method merely requires GWAS summary statistics and an “LD score” (the amount of genetic variation tagged by a SNP) reference panel.

Bivariate LD Score regression utilizes the following moment condition:

$$(6) \quad E[z_{1j}z_{2j}] = \text{Intercept} + \frac{\sqrt{N_1N_2}}{M} \text{Cov}_g \ell_j,$$

where z_{kj} is the z -statistic of SNP j from the GWAS of trait k ($k = 1, 2$), *Intercept* is the regression intercept, N_k is the sample size of the GWAS of trait k , M is the number of SNPs included in the GWAS, Cov_g is the genetic covariance between traits 1 and 2, and ℓ_j is the LD score of SNP j . The slope parameter from a regression of $\hat{z}_{1j}\hat{z}_{2j}$ on $\sqrt{N_1N_2}\ell_j$ can therefore be used to estimate the genetic covariance between the two traits. From separate, univariate LD Score regressions of traits 1 and 2, we can also back out estimates of the respective heritabilities of the two traits, h_{g1}^2 and h_{g2}^2 , and obtain an estimate of the genetic correlation as follows:

$$(7) \quad \hat{r}_g = \frac{\widehat{\text{Cov}}_g}{\sqrt{\hat{h}_{g1}^2 \hat{h}_{g2}^2}}.$$

We used the scores computed by Finucane *et al.*⁹⁴, which use genotypic data from the European-ancestry samples in the 1000 Genomes Project and only HapMap3 SNPs (eur_w_ld_chr, see <https://github.com/bulik/ldsc/wiki/Genetic-Correlation>, accessed on March 14, 2016). As is common in the literature, we restrict our analyses to SNPs with MAF > 0.01; this guarantees all analyses are performed using a set of SNPs that are imputed with reasonable accuracy across all contributing cohorts. The standard errors are estimated by the LDSC software using a block jackknife over the SNPs.

7.2 Pre-selected phenotypes

For our bivariate LD Score analyses, we considered a wide range of phenotypes. First, we considered our supplementary GWAS phenotypes. These include adventurousness, our four main

risky behaviors (automobile speeding propensity, drinks per week, ever smoker, and number of sexual partners) and their first PC (see **Supplementary Table 7.1A1**). We also analyzed additional risky behavior phenotypes for which we ran additional GWAS with only the *first* release of UKB data (for details regarding the methodology and phenotype definitions, see the Appendix at the end of this section); these include age first had sexual intercourse, teenage conception (females only), and use of sun protection (see **Supplementary Table 7.1A2**). Further, we analyzed additional risky behaviors for which we were able to obtain summary statistics from previously published well-powered GWAS. We include the risky behaviors age tobacco smoking onset (among ever-smokers)³⁵ cigarettes per day (among ever-smokers)³⁵, former tobacco smoker (among ever-smokers)³⁵ lifetime cannabis use¹⁰⁰, and self-employed¹⁰¹ (**Supplementary Table 7.1A3**).

Second, we include the *cognition phenotypes* cognitive performance¹⁰², educational attainment¹⁶, and intracranial volume¹⁰³ (**Supplementary Table 7.1B**). Third, we selected the *anthropometric phenotypes* BMI¹⁰⁴ and height¹⁰⁵ (**Supplementary Table 7.1C**). Fourth, we analyzed the *neuropsychiatric phenotypes* ADHD⁶⁹, Alzheimer's disease¹⁰⁶, anxiety disorders¹⁰⁷, autism spectrum disorder¹⁰⁸, bipolar disorder¹⁰⁹, depressive symptoms¹⁸, and schizophrenia⁵⁸ (**Supplementary Table 7.1D**).

Fifth, we analyzed the five *personality phenotypes* that make up the five-factor personality model (agreeableness, conscientiousness, extraversion, neuroticism, and openness to experience) using GWAS summary statistics provided by 23andMe and previously analyzed by Lo *et al.* (2016)¹¹⁰ (**Supplementary Table 7.1E**). Also known as the Big Five, these traits constitute the most widely used taxonomy of personality traits in psychology. The Big Five have roots in Allport & Odbert's lexical hypothesis¹¹¹, which states that individual differences are encoded in language¹¹². This is in contrast to economic preferences such as risk aversion and delay discounting, which are measures of individual heterogeneity that arise in a utility maximization framework. As our sixth *personality phenotype*, we analyzed delay discounting, a measure of impatience, or of the extent to which an individual devalues rewards that are delayed¹¹³.

Recent work in economics has highlighted the importance of personality for economic outcomes, particularly in education, crime, health, and the labor market¹¹⁴. However, little is known about the relationship between the Big Five and economic preferences. In the most comprehensive study of its kind to date, Becker *et al.* (2012)³ find highly significant positive correlations for self-reported risk aversion with openness (0.28) and extraversion (0.26), and negative correlations with conscientiousness (-0.04), agreeableness (-0.14), and neuroticism (-0.09)³.

We also ran GWAS using the first release of UKB data for the *socioeconomic phenotypes* household income and Townsend deprivation index (see **Supplementary Table 7.1E**; for details regarding the methodology and phenotype definitions, see again the Appendix at the end of this section). Finally, we evaluated the genetic correlation between risk and longevity¹¹⁵.

7.3 Results: Genetic correlation between general risk tolerance and pre-selected phenotypes

To estimate the genetic correlations between general risk tolerance and each of the above phenotypes, we used the summary statistics from the meta-analysis of the discovery and replication GWAS. The estimates of the genetic correlations are shown in **Supplementary Table 7.1** and in **Fig. 2**.

7.3.1 *Adventurousness and risky behaviors*

First, we examine the genetic correlation between adventurousness and general risk tolerance. Then, we return to our other supplementary GWAS phenotypes and their genetic correlations with general risk tolerance. These results are reported in **Supplementary Table 7.1A1**. As expected, the estimated genetic correlation with adventurousness is high and statistically significant ($\hat{r}_g = 0.834$, $SE = 0.011$). The correlation with automobile speeding propensity is moderately high and highly statistically significant ($\hat{r}_g = 0.448$, $SE = 0.021$), consistent with individuals with more risk-tolerance increasing alleles being more likely to exceed the automobile speed limit. For drinks per week the genetic correlation is positive and highly statistically significant ($\hat{r}_g = 0.254$, $SE = 0.021$), and the genetic correlation is comparable for ever smoker ($\hat{r}_g = 0.246$, $SE = 0.022$), implying that higher risk tolerance is genetically associated with more risky health behaviors. The genetic correlation estimates are highly significant for number of sexual partners ($\hat{r}_g = 0.520$, $SE = 0.019$). Finally, the genetic correlation for the first PC of these four risky behaviors, which can be interpreted as a general factor of risky behavior, is interestingly both moderately high in absolute value and highly statistically significant ($\hat{r}_g = 0.500$, $SE = 0.018$). Thus, all of our supplementary GWAS phenotypes are significantly genetically correlated in the expected direction with the primary general-risk-tolerance phenotype.

Next, let us turn to the three additional UKB risky behaviors (for which we ran GWAS using only the first release of UKB data) and their genetic correlations with general risk tolerance. The results are reported in **Supplementary Table 7.1A2**. The genetic correlation estimates are highly significant for age first had sexual intercourse ($\hat{r}_g = -0.332$, $SE = 0.032$) and teenage conception ($\hat{r}_g = 0.246$, $SE = 0.049$). Together with our results of number of sexual partners, this implies that higher risk tolerance is genetically associated with more risky sexual behavior; this is also consistent with a finding from a recent GWAS of age at first sexual intercourse⁶⁶. Use of sun protection has an insignificant genetic correlation with general risk tolerance, and we suspect that this phenotype may be more highly correlated with skin pigmentation and predisposition to skin cancer than to general risk tolerance.

All seven of our statistically significant genetic correlation estimates for the aforementioned risky behaviors have signs that are in the direction that would be expected based on the corresponding phenotypic correlations, and the absolute values of the estimates are higher than those of these phenotypic correlations.

Lastly, **Supplementary Table 7.1A3** reports the estimates for the risky behaviors for which we were able to obtain summary statistics from previously published, well-powered GWAS. One of the three cigarette-related phenotypes is moderately and significantly genetically correlated with general risk tolerance: former tobacco smoker (among ever-smokers) ($\hat{r}_g = -0.131$, $SE = 0.055$). However, age of tobacco smoking onset (among ever-smokers) and cigarettes per day (among ever-smokers) are not significantly genetically correlated with general risk tolerance. These results suggest that, while higher risk tolerance may be genetically associated with some risk-related smoking behaviors such as smoking initiation (as suggested by the significant genetic correlation with ever smoker) and cessation, risk tolerance may not necessarily be genetically correlated with smoking addiction (as captured by cigarettes per day).

For lifetime cannabis use, the genetic correlation is positive and highly statistically significant ($\hat{r}_g = 0.313$, $SE = 0.057$), implying that risk tolerance is genetically associated with risk-seeking

cannabis use behavior. The genetic correlation estimate for self-employed is significant and large in magnitude ($\hat{r}_g = 0.672$, $SE = 0.259$), implying that higher risk tolerance is genetically associated with a common proxy for entrepreneurship. The point estimate for this phenotype is the highest among any of the phenotypes tested. However, in interpreting this correlation it is important to note that the LD Score regression heritability of the self-employment phenotype is low and not significantly different from zero: 0.0126 ($SE = 0.0106$). Indeed, the standard errors on the estimate are quite large ($SE = 0.259$). This might be due to the fairly small sample size of the self-employment GWAS ($n = 50,627$).

All 11 of our statistically significant genetic correlation estimates for the risky behaviors have signs that imply higher risk tolerance is associated with riskier behaviors.

7.3.2 *Cognition phenotypes*

Two of the three cognition phenotypes are significantly, though weakly, genetically correlated with general risk tolerance: educational attainment ($\hat{r}_g = 0.099$, $SE = 0.022$) and intracranial volume ($\hat{r}_g = 0.144$, $SE = 0.059$). These results are consistent with the well-established positive correlation between risk preferences and educational attainment in the literature². The absence of genetic correlation between risk tolerance and cognitive performance ($\hat{r}_g = 0.012$, $SE = 0.063$) is surprising, given that risk tolerance and cognitive performance have been shown to be positively correlated at the phenotypic level^{116,117}. A possible explanation is that cognitive performance tends to correlate positively with avoidance of harmful risky situations but negatively with avoidance of beneficial risky situations¹¹⁸, and that our measure of risk tolerance captures behavior in both types of situations in such a way that the effects cancel out when estimating the genetic correlation.

7.3.3 *Anthropometric phenotypes*

Height is not significantly genetically correlated with general risk tolerance. BMI, on the other hand, is significantly, albeit weakly, positively correlated ($\hat{r}_g = 0.053$, $SE = 0.021$).

7.3.4 *Neuropsychiatric phenotypes*

Among the neuropsychiatric traits, we find moderate (and highly significant) genetic correlations with ADHD ($\hat{r}_g = 0.247$, $SE = 0.033$), anxiety disorders, ($\hat{r}_g = 0.214$, $SE = 0.089$), autism spectrum disorder ($\hat{r}_g = -0.105$, $SE = 0.050$), bipolar disorder ($\hat{r}_g = 0.214$, $SE = 0.035$) and schizophrenia ($\hat{r}_g = 0.173$, $SE = 0.021$). The positive and significant genetic correlation with ADHD is consistent with the significant effect of our polygenic score of general risk tolerance on ADHD in an independent sample (**Supplementary Information section 10**). We do not, however, find significant genetic correlations with either Alzheimer's disease or depressive symptoms.

7.3.5 *Personality phenotypes*

Agreeableness and conscientiousness are not significantly correlated with general risk tolerance. However, the genetic correlations between general risk tolerance and extraversion ($\hat{r}_g = 0.505$, $SE = 0.027$), neuroticism ($\hat{r}_g = -0.420$, $SE = 0.038$), and openness ($\hat{r}_g = 0.332$, $SE = 0.031$), are moderately high in absolute value and highly significant. The direction of the genetic correlation between general risk tolerance and all five personality traits is in line with the literature on the

phenotypic correlations of these traits outlined above³. Our results for extraversion, neuroticism, and openness are also in line with the signs of the significant coefficients on the polygenic score for general risk tolerance in regressions of each of these personality phenotypes on the score, as highlighted in **Supplementary Information section 10** (although our predictive power for these traits is quite low).

Interestingly, we find a significant and moderate positive genetic correlation between general risk tolerance and delay discounting ($\hat{r}_g = 0.210$, $SE = 0.101$), indicating that higher risk tolerance is associated with higher impatience. Our estimate of this genetic correlation is similar to our estimate of the genetic correlation between general risk tolerance and ADHD; Sanchez-Roige *et al.*¹¹³ note that delay discounting may act as an endophenotype for ADHD, and the similarity in the two genetic correlation estimates support this conclusion.

7.3.6 *Socioeconomic phenotypes and longevity*

The genetic correlation estimates for both household income ($\hat{r}_g = 0.215$, $SE = 0.033$) and for Townsend score ($\hat{r}_g = 0.185$, $SE = 0.047$) are positive and significant, implying that higher risk tolerance is genetically associated with higher earnings and social deprivation. The genetic correlation with Townsend score is higher than even the phenotypic correlation. We do not find a significant genetic correlation with longevity.

7.3.7 *Summary of Findings*

In sum, general risk tolerance tends to be genetically correlated with adventurousness and with the risky behaviors involving automobile speeding propensity, substance use, sexual activity, and self-employment. Importantly, our estimates have signs that are consistent with higher self-reported risk tolerance being associated with riskier behavior. General risk tolerance is also genetically correlated with the neuropsychiatric phenotypes ADHD, anxiety disorders, autism spectrum disorder, bipolar disorder, and schizophrenia, but not with Alzheimer's disease or depressive symptoms. General risk tolerance is also correlated with the Big Five personality phenotypes extraversion, neuroticism, and openness, but not with agreeableness or conscientiousness, and is correlated with delay discounting.

Our results also point toward distinctions in the genetic correlation between general risk tolerance and externalizing and internalizing behaviors and disorders. Externalizing behaviors and disorders are those in which individuals tend to express maladaptive thoughts and feelings toward others or their environment, while internalizing behaviors and disorders are those in which individuals tend to express thoughts and feelings inward. Overall, we find significant genetic correlations with behaviors and disorders typically classified as externalizing¹¹⁹, such as addiction (smoking cigarettes, cannabis use, and drinking) and ADHD as well as thought disorders related to externalizing disorders such as bipolar disorder and schizophrenia. Conversely, we find little evidence of significant genetic correlation with internalizing behaviors or disorders, such as depression. We also find evidence confirming the well-documented phenotypic correlation between risk taking and cognitive performance and educational attainment (e.g., in the domain of financial risk taking^{2,4,116}).

7.4 Other results

7.4.1 Genetic correlation between males and females

Phenotypically, more than 34% of males in the UKB are categorized as risk tolerant (based on their answers to the general-risk-tolerance question), whereas only approximately 19% of females are categorized as risk tolerant. To test the extent to which the genetics of general risk tolerance differs between males and females, we conducted GWAS of general risk tolerance separately for males and females. We followed the same methodology and QC protocol for these two sex-specific GWAS as for our other GWAS in the full release of UKB data. **Supplementary Table 7.2** reports the SNP-based heritabilities and other relevant summary statistics from the LD Score regressions. General risk tolerance is slightly more heritable for males ($h^2 = 0.064$, $SE = 0.004$) than for females ($h^2 = 0.055$, $SE = 0.003$). We then used bivariate LD Score regression to calculate the genetic correlation between these two samples, and obtained an estimate of $\hat{r}_g = 0.822$ ($SE = 0.033$). The genetic correlation is smaller than unity, pointing to some heterogeneity across females and males, but high enough to justify our approach of pooling males and females in our other analyses to maximize statistical power (see section 2 of the Supplementary Note of Okbay *et al.* (2016) for derivations the demonstrate this).

7.4.2 Genetic correlation between the discovery and replication GWAS and between the 23andMe and UKB cohorts

We estimated the genetic correlation between our discovery and our replication GWAS of general risk tolerance. We find a high genetic correlation that cannot be statistically distinguished from unity ($\hat{r}_g = 0.8344$, $SE = 0.1289$), suggesting the genetic underpinnings of general risk tolerance do not vary much between our discovery and replication samples.

As discussed in **Supplementary Information sections 1 and 3.3**, we also estimated the genetic correlation between: (1) the UK Biobank general-risk-tolerance GWAS and the 23andMe general-risk-tolerance GWAS; (2) the UK Biobank general-risk-tolerance GWAS and the replication GWAS; (3) the 23andMe general-risk-tolerance GWAS and the replication GWAS; and (4) the discovery and replication GWAS of general risk tolerance. For (1) we find a moderately high, positive genetic correlation ($\hat{r}_g = 0.767$, $SE = 0.021$) that is distinguishable from unity. For (2) we find a moderately high, positive genetic correlation ($\hat{r}_g = 0.828$, $SE = 0.135$); for (3) we find ($\hat{r}_g = 0.759$, $SE = 0.126$). Finally, for the final correlation (4), we find ($\hat{r}_g = 0.834$, $SE = 0.129$). The last three correlations are all indistinguishable from unity.

7.4.3 Relationship between the genetic correlation and the fraction of overlapping lead loci across pairs of phenotypes

Following a suggestion by a Referee, we conducted an exploratory analysis of the relationship between the genetic correlation between a pair of phenotypes and a measure of the fraction of lead SNPs (and SNPs in weak LD, $r^2 > 0.1$) that overlap across the phenotypes' GWAS. We examined this relationship across six pairs of phenotypes, each comprising general risk tolerance and one of the six supplementary phenotypes. Our measure of the fraction of lead SNPs that overlap across phenotypes 1 and 2 is $\frac{q_{12}}{\sqrt{q_1 q_2}}$, where q_{12} is the number of overlapping lead SNPs and q_i is the number of lead SNPs for phenotype i ($i = 1, 2$). The number of overlapping lead SNPs is defined as the

number of lead SNPs of the supplementary phenotype that are in weak LD ($r^2 > 0.1$) with a general-risk-tolerance lead SNP. We estimated a positive correlation of 0.34 between our measure of the fraction of overlapping lead SNPs and the genetic correlation, consistent with the intuition that pairs of phenotypes with higher genetic correlations will tend to have more overlapping lead SNPs. However, this estimate is not statistically significant ($P = 0.51$), which is not surprising given how few data points were used in the analysis, which limited statistical power.

7.5 Comparison of the genetic and phenotypic correlations

In **Supplementary Table 1.3** we show phenotypic correlations for our primary GWAS phenotype, general risk tolerance, and for our five supplementary phenotypes whose GWAS included data from the UKB: automobile speeding propensity, drinks per week, ever smoker, number of sexual partners, and the first PC of these four risky behaviors (the GWAS of our other supplementary phenotype, adventurousness, used 23andMe data only). Panel **A** shows values calculated naively assuming Pearson correlation coefficients for all of our variables using the “`pwcorr`” command in Stata. In the first column, we show test-retest correlations between the first and the second measurements if each phenotype in the UKB, to give a sense of the test-retest reliability of each phenotype. In the subsequent correlation matrix, coefficients below the diagonal are uncorrected Pearson coefficients. Above the diagonal, we adjust the correlation estimate for each pair of variables for measurement error by dividing it by the square root of the product of the two test-retest correlations for the two variables^{ee}.

In Panel **B**, we recalculate Panel **A**, but this time we use the polychoric package in Stata to allow the correct estimation of tetrachoric (between two ordinal variables) or polyserial correlations (between an ordinal and a continuous variable). Again, uncorrected correlation coefficients are below the diagonal, while correlations corrected for measurement error are above the diagonal.

In summary, both correcting for measurement error (Panel **A** and **B**, above the diagonals) and specifying the correct type of correlation coefficient (Panel **B**) raises the value of the correlation estimates, so that our highest estimates are in Panel **B**, above the diagonal. While some correlations are quite high (especially between the first PC phenotype and the other phenotypes), the others remain relatively low, including for the correlations between general risk tolerance and the five supplementary phenotypes.

In **Supplementary Table 7.1**, Panel **A1**, we compare the *genetic* correlations with general risk tolerance to the *phenotypic* correlations with general risk tolerance. Even after specifying the correct type of correlation (Pearson or polyserial) and adjusting for measurement error, most genetic correlations remain considerably higher than the phenotypic correlations.

These results are relevant to the ongoing debate about the extent to which risk tolerance is a “domain-general” versus “domain-specific” trait. Low phenotypic correlations among measures of risky behaviors in various domains have led some researchers to conclude that risk tolerance is highly domain-specific^{120,121}. The comparatively large genetic correlations we estimate support the view that a general factor of risk tolerance partly accounts for cross-domain variation in risky behavior^{6,122} and imply that this factor is genetically influenced, while the lower phenotypic

^{ee} This is a standard method to correct for measurement error attenuation in the correlation estimates (see, e.g., https://en.wikipedia.org/wiki/Correction_for_attenuation).

correlations suggest that environmental factors are more important contributors to domain-specific behavior.

7.6 Appendix: GWAS of other risky behaviors and of socioeconomic phenotypes using the first release of UKB data

While we used existing published GWAS results for most phenotypes, we conducted our own GWAS using the full release of UKB data for our four main risky behaviors and their first PC. We describe the coding of our four main risky behaviors and their first PC in **Supplementary Information section 1.2** (for phenotypic correlations of these five behaviors see **Supplementary Table 1.3**). We also conducted our own GWAS using only the first release of UKB data for five phenotypes: age first had sexual intercourse^{ff} ($n = 98,956$), teenage conception among females ($n = 40,077$), use of sun protection, household income ($n = 97,059$), and Townsend deprivation index^{gg} score ($n = 112,192$). Throughout these analyses we dropped participants who answered “Do not know” or “Prefer not to answer” and averaged data across two assessment visits when possible.

The five remaining UKB phenotypes for which we ran GWAS in only the first release of UKB data were coded as follows:

- Age first had sexual intercourse: UKB respondents were asked “What was your age when you first had sexual intercourse? (Sexual intercourse includes vaginal, oral or anal intercourse)?” We dropped anyone reporting “Never had sex” or an age of first sexual encounter at less than 12 (given the high likelihood of associated abuse or misreporting). We then normalized the measure separately for males and females.
- Teenage conception among females: UKB females who bore at least one child were asked “How old were you when you had your FIRST child?” We recoded this variable into a case-control binary variable. Females reporting age of first live birth between 13 and 20 (inclusive) were coded as cases ($n = 6,285$), while females reporting higher ages of first live birth were coded as controls ($n = 33,792$). Childless females were dropped.
- Use of sun protection: UKB respondents were asked “Do you wear sun protection (e.g. sunscreen lotion, hat) when you spend time outdoors in the summer?”; eligible responses ranged from “1. Never/rarely” to “4. Always,” and also included “5. Do not go out in sunshine.” We dropped all participants who answered “5. Do not go out in sunshine” and normalized the resulting categorical variable separately for males and females.
- Household income: UKB respondents were asked “What is the average total income before tax received by your HOUSEHOLD?”; eligible responses were “1. Less than £18,000,” “2. £18,000 to £30,999,” “3. £52,000 to £100,000,” “5. Greater than £100,000.” We normalized the resulting categorical variable.
- Townsend deprivation index score: This score measures local social deprivation based on the preceding national census output areas; a higher score implies more social deprivation. Each participant was assigned a score corresponding to the output area in which their

^{ff} Day *et al.*⁶⁶ report results from a GWAS of age of first sexual encounter. We conduct similar analyses ourselves here. We treat the phenotype slightly differently by separately normalizing phenotypes among males and females, and then conducting our GWAS on the combined sample.

^{gg} Hill *et al.*²⁷⁰ report results from GWAS for household income and the Townsend deprivation index in the UKB. We ran our own analyses because we could not find the summary statistics from their GWAS in the public domain.

postcode is located. The score is calculated by the UKB immediately prior to each participant joining the dataset. We normalized the scores for our analysis.

We followed the same methodology and QC protocol for these five additional GWAS as for our discovery GWAS of general risk tolerance and our GWAS of the four main risky behaviors and their first PC in the full release of UKB data (see **Supplementary Information section 2**), except that (1) we only used unrelated individuals in the first release of UKB data; (2) we conducted the association analyses with SNPtest v.2.4.1⁹² (instead of BOLT-LMM⁴⁰); (3) the summary statistics were quality controlled using the 1000 Genomes phase 3 reference panel; and (4) we only used SNPs with MAF > 0.005 instead of MAF > 0.001 (the latter does not affect the analyses reported in this section, as we restrict these analyses to SNPs with MAF > 0.01). **Supplementary Table 7.2B** reports the various statistics outputted by LD Score regressions for each of these five additional GWAS. The third column shows estimates of the SNP-based heritabilities. All five phenotypes have a higher SNP-based heritability than general risk tolerance, except Townsend score. The sexual activity phenotypes exhibit particularly high heritabilities. Age first had sexual intercourse has the highest SNP-based heritability: 0.167 ($SE = 0.009$). The only estimate of the LD Score regression intercept that is significantly different from one is for household income, for which the intercept is 1.035 ($SE = 0.008$). By comparison, the mean χ^2 statistics for the SNPs in the LD Score regressions are larger than 1.10 for three of the five GWAS, including household income for which the mean χ^2 statistic is 1.198 (the exceptions are teenage conception and Townsend score, for which the mean χ^2 statistics are 1.074 and 1.094, respectively). These estimates imply that only a small part of the observed inflation in the mean χ^2 statistics of the GWAS is likely to be accounted for by confounding bias (due to population stratification, cryptic relatedness, or other confounds), rather than by polygenic signals. Additional details for these five additional GWAS are available upon request.

8 Proxy-phenotype analyses

8.1 Introduction

We conducted proxy-phenotype analyses¹⁰² to search for additional SNPs that affect phenotypes that are plausibly related to general risk tolerance. These analyses allow us to test whether SNPs that are strongly associated with a “first-stage” phenotype are enriched for association with a related “second-stage” phenotype, and potentially to identify new SNPs associated with the second-stage phenotype. Proxy-phenotype analyses leverage the fact that, if the first- and second-stage phenotypes are genetically correlated, SNPs associated with the first-stage phenotype should have a higher probability of being associated with the second-stage phenotype, compared to what we would expect by chance.

In our study, the first-stage phenotype is always general risk tolerance, and the first-stage analysis was conducted in our discovery GWAS. We consider eight primary second-stage phenotypes: age of smoking onset, cigarettes per day (CPD), smoking cessation (which compares former smokers to current smokers), lifetime cannabis use, self-employment status (an indicator of whether an individual is self-employed or not), ADHD, bipolar disorder, and schizophrenia. We chose these second-stage phenotypes for the following two reasons. First, summary statistics from GWAS of these phenotypes that did not include the UKB are publicly available or could be obtained. Second, these phenotypes are plausibly related to general risk tolerance: the first five phenotypes are risky behaviors and all eight are at least moderately genetically correlated with general risk tolerance (**Supplementary Information section 7.3**). We also used height as a negative control.

8.2 Methodology

8.2.1 Data and Setup

We used our discovery GWAS of general risk tolerance as our first-stage analysis. The second-stage lookups were all performed using summary statistics from previous GWAS. We obtained the summary statistics from the following sources: The Tobacco and Genetics Consortium (2010)³⁵ for the three tobacco smoking phenotypes, the International Cannabis Consortium lifetime for cannabis use (Stringer *et al.* 2016)¹⁰⁰, the Entrepreneur Consortium for self-employment (van der Loos *et al.* 2013)¹⁰¹, the Psychiatric Genomics Consortium for ADHD, bipolar disorder, and schizophrenia (Demontis *et al.* 2017, Sklar *et al.* 2011, and Ripke *et al.* 2014, respectively)^{58,69,109}, and the GIANT consortium for height (Wood *et al.* 2014)¹⁰⁵. Our methodology follows Okbay *et al.* (2016)¹⁸ and involves two main stages.

8.2.2 Stage 1: Constructing the set of lead and proxy-lead SNPs

The first stage involves constructing the set of lead SNPs from the first-stage analysis. Unlike Okbay *et al.*, we only used the lead SNPs (i.e., the approximately independent SNPs with P values $< 5 \times 10^{-8}$) from our discovery GWAS of general risk tolerance as candidate SNPs for the second stage^{hh}.

For brevity, we illustrate the steps involved for cigarettes per day (CPD), but analogous procedures apply for each of the other second-stage phenotypes. Our discovery GWAS of general risk tolerance identified 124 lead SNPs. Of these, 52 SNPs were directly available in the CPD summary statistics, whereas 72 were either not available or their GWAS sample sizes were too small to meet

^{hh} Okbay *et al.* used all SNPs with a P value less than 1×10^{-4} , rather than just the lead SNPs.

our inclusion criterion: to ensure the quality of the estimates for the second-stage SNPs, we limit our lookup procedure to second-stage SNPs with GWAS sample sizes of at least one-half of the maximum sample size for that GWAS. For each of these 72 SNPs, we determined whether there exists a suitable “proxy-lead SNP” that satisfies three conditions: (1) the SNP is in high LD ($r^2 > 0.8$) with the risk-tolerance associated SNP; (2) the SNP is available in the summary statistics of both the discovery GWAS of general risk tolerance and the GWAS of CPD; and (3) the SNP has a CPD sample size of at least one-half the maximum sample size in the CPD GWAS. To determine the set of SNPs in high LD with the risk-associated SNP, we employ PLINK⁴³ with our main reference panelⁱⁱ. A proxy-lead SNP was available for an additional 44 of the 72 SNPs (with r^2 's ranging from 0.80 to 1.00, with a mean $r^2 = 0.94$). Whenever more than one proxy was available for a SNP, we chose the proxy with the largest r^2 . Ties were broken by choosing the closest SNP to the original lead SNP in base pair distance. Our final list of lead and proxy-lead SNPs for CPD therefore contained $52 + 44 = 96$ SNPs.

8.2.3 *Stage 2, part 1: SNP lookup and search for previously identified significant loci*

We *individually* tested each of the k lead and proxy-lead SNPs for experiment-wide significance by examining whether each is significantly associated with the second-stage phenotype, using a Bonferroni-corrected significance level of $0.05/k$ (for CPD, $k = 96$). We refer to the lead and proxy-lead SNPs that reached Bonferroni-corrected significance in the second stage as “second-stage hits.”

For each second-stage hit, we then identified the set of SNPs in weak LD with the second-stage hits by identifying the SNPs in a 1,000 kb window around the corresponding first-stage lead SNP and with $r^2 > 0.1$ with that corresponding first-stage lead SNP (we used PLINK⁴³ and our main reference panel to compute LD). We then checked if any of these SNPs are genome-wide significant in the second-stage summary statistics. This allowed us to determine whether the second-stage hits tag genomic regions that were previously found to be genome-wide significant in the GWAS of the second-stage phenotype.

8.2.4 *Stage 2, part 2: Testing the set of lead and proxy-lead SNPs for enrichment and sign concordance*

For each second-stage phenotype, we performed a non-parametric Mann-Whitney test⁸⁵ of joint enrichment to test whether the set of lead and proxy-lead SNPs have a P value distribution that is significantly different from the P value distribution of a randomly-chosen, matched set of “comparison SNPs” in the second-stage summary statistics. Because we expect the second-stage phenotypes to be highly polygenic, with many SNPs having weak but true associations, it would have been inappropriate to test the null hypothesis that the P value distribution of the lead and proxy-lead SNPs is uniform. For each lead and proxy-lead SNP, the comparison SNPs are randomly selected from among the set of SNPs that have a minor allele frequency within one percentage point of the lead or proxy-lead SNPⁱⁱ. As in Okbay *et al.* (2016)¹⁸, we generated 1,000 comparison SNPs for each of the k lead or proxy-lead SNPs, and compared the P value distribution for this group of $k \times 1000$ SNPs with that of the k lead or proxy-lead SNPs.

We also conducted a sign test to assess whether the lead and proxy-lead SNPs have effects in the predicted (or concordant) direction in the second stage. For cigarettes per day (CPD), lifetime

ⁱⁱ Throughout this section, we employ our main reference panel to compute the LD between SNPs.

^{jj} The matched SNPs are drawn with replacement from the set of SNPs in the second-stage summary statistics that excludes the Y lead or proxy-lead SNPs.

cannabis use, self-employment status, ADHD, bipolar disorder, and schizophrenia (for which a higher phenotype value corresponds to more risk taking, and which we estimated to be positively genetically correlated with general risk tolerance) we classified a SNP as having an effect “in the predicted direction” if the sign of its effect is concordant with that for general risk tolerance. For age of smoking onset and smoking cessation (for which a higher phenotype value corresponds to less risk taking, and which we estimated to be negatively genetically correlated with general risk tolerance) we classified a SNP as having an effect “in the predicted direction” if the sign of its effect is discordant with that for general risk tolerance. For example, a SNP that increases general risk tolerance and decreases the age of smoking onset has an effect in the predicted direction. For ADHD, bipolar disorder, schizophrenia, and height, we did not predict the direction of the SNPs’ effects; we simply conducted a sign test to establish whether SNPs’ effects are more likely to be sign concordant with those for general risk tolerance than expected by chance.

8.3 Results from proxy-phenotype enrichment analyses

Q-Q plots for the lead and proxy-lead SNPs in each second-stage phenotype are shown in **Extended Data Fig. 8.1**. These plots show results from the sign test mentioned above and highlight any second-stage hits. Further details on each of the lead and proxy-lead SNPs are reported in **Supplementary Table 8.1**, which reports summary statistics from the first and second stages for the lead SNPs associated with general risk tolerance and their proxies.

8.3.1 *Do risk-tolerance-associated SNPs predict smoking behaviors?*

For CPD, 52 lead SNPs are directly available in the second-stage summary statistics, and we identified 44 proxy-lead SNPs. Out of 96 lead or proxy-lead SNPs, 51 (53.1%) have signs in the predicted direction ($P = 0.31$). There are no second-stage hits. Moreover, the Mann-Whitney test of joint enrichment fails to reject the null hypothesis that the P values of the lead and proxy-lead SNPs are drawn from the same P value distribution as a set of randomly-selected SNPs ($P = 0.88$).

For age of smoking onset, 52 lead SNPs are directly available in the second-stage summary statistics, and we identified 45 proxy-lead SNPs. Of a total of 97 lead or proxy-lead SNPs, 54 (55.7%) have signs in the predicted direction ($P = 0.15$). There were again no second-stage hits, and the Mann-Whitney test of joint enrichment fails to reject the null ($P = 0.48$).

For smoking cessation, 53 lead SNPs are directly available in the second-stage data, and we identified 44 proxy-lead SNPs. Of a total of 97 lead or proxy-lead SNPs, 61 (62.9%) have signs in the predicted direction ($P = 0.007$). There were no second-stage hits. The Mann-Whitney test of joint enrichment fails to reject the null ($P = 0.11$).

8.3.2 *Do risk-tolerance-associated SNPs predict lifetime cannabis use?*

For lifetime cannabis use, 117 lead SNPs are directly available in the second-stage data, with no additional proxy-lead SNPs. Of these lead SNPs, 76 out of 117 (65.0%) have signs in the predicted direction ($P = 8 \times 10^{-4}$). One SNP, rs993137, reaches Bonferroni-corrected significance ($P = 1.7 \times 10^{-4}$, before Bonferroni correction) and is thus a second-stage hit, and the sign of the effect of this SNP is in the predicted direction. This SNP does not tag any previously identified genome-wide hit for lifetime cannabis use, and is thus a novel association. The Mann-Whitney test of joint enrichment fails to reject the null hypothesis ($P = 0.13$).

Interestingly, rs993137 falls within the *CADM2* gene, which was found to be significantly associated with lifetime cannabis use in a gene-based test by Stringer *et al.* (2016)¹⁰⁰. Stringer *et*

al. also note that *CADM2* has previously been associated with body mass index (BMI), processing speed, and autism disorders; these phenotypes themselves have been previously associated with cannabis use^{123–125}. It is noteworthy that *CADM2* contains lead SNPs for all our primary and supplementary GWAS. The NHGRI-EBI GWAS Catalog database⁶⁵ reports previous associations with age at menarche, cognitive function, and educational attainment, among other phenotypes, as well as those mentioned by Stringer *et al.* (2016).

8.3.3 Do risk-tolerance-associated SNPs predict self-employment?

For self-employment, 56 lead SNPs are directly available in the second-stage data, along with 43 proxy-lead SNPs. Of a total of 99 lead or proxy-lead SNPs, 64 (64.6%) have signs in the predicted direction ($P = 0.002$). There is one second-stage hit, SNP rs7387531 ($P = 1.1 \times 10^{-4}$, before Bonferroni correction), and the sign of its effect is in the predicted direction.^{kk, ll} To our knowledge, no robust association has previously been reported between a genetic variant and self-employment; thus, if the association with rs7387531 is robust, this would be the first genetic variant to be found to be significantly associated with self-employment.

Lastly, the Mann-Whitney test of joint enrichment fails to reject the null hypothesis ($P = 0.49$). The significant SNP rs7387531 is located within a candidate inversion on chromosome 8 (~7.89 to 11.79 Mb), and that candidate inversion contains lead SNPs for our GWAS of general risk tolerance, adventurousness, automobile speeding propensity, ever smoker, and number of sexual partners. The NHGRI-EBI GWAS Catalog database⁶⁵ reports SNP associations within the breakpoints of the candidate inversion with many other phenotypes, among them are neuroticism, extraversion, schizophrenia, and chronotype.

8.3.4 Do risk-tolerance-associated SNPs predict ADHD?

For ADHD, 116 lead SNPs are directly available in the second-stage data, with one additional proxy-lead SNP. Of a total of 117 lead or proxy-lead SNPs, 74 (63.2%) SNPs have concordant signs ($P = 0.003$). There are four second-stage hits (rs10905461, $P = 5.0 \times 10^{-5}$ before Bonferroni correction; rs3764002, $P = 2.4 \times 10^{-4}$; rs7783012, $P = 4.4 \times 10^{-6}$; and rs786250, $P = 2.6 \times 10^{-5}$) and all of these have concordant signs. These SNPs are all in new loci: none of them tag any previously identified genome-wide associations for ADHD. The Mann-Whitney test of joint enrichment rejects the null hypothesis of no-enrichment ($P = 0.008$).

None of the four Bonferroni-significant SNPs are located within any of the long-range LD regions and candidate inversions in which we found lead SNPs for all or most of our primary and supplementary GWAS (**Supplementary Information section 3.2**).

We also note that ADHD is significantly associated with our polygenic score of general risk tolerance (**Supplementary Information section 10**) and, as with all second-stage traits we analyze here (except height), it is also significantly genetically correlated with general risk tolerance.

^{kk} In an *ex post* analysis, we looked up rs7387531 in the summary statistics of the replication GWAS of self-employment from van der Loos *et al.* (2013)¹⁰¹. rs7387531's association with self-employment did not replicate ($P = 0.061$ on a two-sided test, but with the wrong sign). However, this replication attempt was severely underpowered: the replication sample was small, comprising only the STR cohort ($n = 3,271$). Further, rs7387531 had an R^2 of ~0.004% in the discovery GWAS of general risk tolerance; due to the winner's curse, the true R^2 is likely to be smaller than that, and the R^2 of rs7387531 on self-employment could be even smaller.

^{ll} We assume that higher general risk tolerance leads to a higher probability of being self-employed. For the opposite view, see ref.²⁷¹.

8.3.5 *Do risk-tolerance-associated SNPs predict bipolar disorder?*

For bipolar disorder, 47 lead SNPs are directly available in the second-stage data, with 49 additional proxy-lead SNPs. Of a total of 96 lead or proxy-lead SNPs, 64 (66.7%) SNPs have concordant signs ($P = 7.1 \times 10^{-4}$). There are no second-stage hits. The Mann-Whitney test of joint enrichment fails to reject the null hypothesis of no-enrichment ($P = 0.87$).

8.3.6 *Do risk-tolerance-associated SNPs predict schizophrenia?*

For schizophrenia, 122 lead SNPs are directly available in the second-stage data, with no additional proxy-lead SNPs. Of a total of 122 lead or proxy-lead SNPs, 85 (70.0%) SNPs have concordant signs ($P = 8.3 \times 10^{-6}$). There are sixteen second-stage hits, and 13 of these have concordant signs. Four of these SNPs do not tag any loci that were previously identified in published GWAS on schizophrenia (rs13327339, $P = 4.8 \times 10^{-5}$ before Bonferroni correction; rs1374197, $P = 4.8 \times 10^{-5}$; rs2357023, $P = 4.0 \times 10^{-4}$; and rs3764002, $P = 3.8 \times 10^{-4}$); two of these four SNPs (rs1374197 and rs2357023) have concordant signs. rs3764002 was also a second-stage hit for ADHD.

Four of the 16 Bonferroni-significant SNPs are located within long-range LD regions or a candidate inversion in which we found lead SNPs in all or most of our primary and supplementary GWAS (**Supplementary Information section 3.2**). The SNP rs3849046 is located within a long-range LD region on chromosome 5 (~135.4 to 138.4 Mb) in which we found two lead SNPs for general risk tolerance, but no lead SNPs for any other main GWAS. The SNP rs1417998 is located in the long-range LD region on chromosome 6 (~25.3 to 33.4 Mb) that covers the HLA-complex, and that region contains lead SNPs for all our GWAS except drinks per week (for which it contains a suggestive association). The SNPs rs624244 and rs1531518 are both located within a candidate inversion on chromosome 18 (~49.1 to 55.5 Mb), and that region is noteworthy because it contains lead SNPs for all our main GWAS.

The Mann-Whitney test of joint enrichment rejects the null hypothesis of no-enrichment ($P = 3.0 \times 10^{-9}$) relative to randomly selected SNPs. Although schizophrenia has been shown to be very polygenic¹²⁶, the Mann-Whitney test rejects the hypothesis that the observed enrichment is due to polygenic inflation of test statistics over the entire genome. However, it is possible that the overlap between risk tolerance and schizophrenia is explained by enrichment of broad classes of SNPs (e.g., functionally important or conserved regions), rather than specific shared pathways. This is because the Mann-Whitney null distribution is only matched on minor allele frequency; however, the null sample could be further matched based on other attributes (e.g., LD score, functional annotations) to examine whether the joint enrichment is accounted for by these attributes.

These 16 second-stage hits for schizophrenia, together with the strong enrichment of the general-risk-tolerance lead SNPs in the schizophrenia GWAS as well as the high sign concordance of their effects on schizophrenia, suggest that part of the genetic signal for schizophrenia and general risk tolerance may be concentrated in the same genomic regions. We also note that five of our general-risk-tolerance lead SNPs are located in loci that had been found by previous GWAS to be associated with schizophrenia (as we detail in **Supplementary Information section 3.3**).

8.3.7 *Do risk-tolerance-associated SNPs predict height?*

For height, 57 lead SNPs are available in the data, along with 42 proxy-lead SNPs. Out of these 99 lead or proxy-lead SNPs, 53 (53.5%) have concordant signs ($P = 0.27$), and eight are second-

stage hits. Four of these eight SNPs have concordant sign, and all tag loci with previously identified genome-wide associations. The Mann-Whitney test of joint enrichment rejects the null hypothesis that the second-stage P values are drawn from a null distribution ($P = 1.4 \times 10^{-5}$) and reveals much enrichment (as can be seen from **Extended Data Fig. 8.1**).

These result for height partially contradicting our initial framing of height as a negative control. Such enrichment in the absence of genetic correlation between two traits could be due to a number of reasons¹²⁷, including an enrichment of all polygenic traits for certain regions in the genome (e.g., functional or evolutionary conserved regions). As discussed above, the Mann-Whitney null distribution does not account for this form of regional polygenic enrichment. We do not further explore this here, but we note that this may be an interesting question for future research.

9 Multi-trait Analysis of GWAS (MTAG)

9.1 Introduction

Because many phenotypes are genetically correlated^{24,87}, information contained in the GWAS of different but related phenotypes can be used to increase detection and the predictive power of our analysis. We used Multi-trait Analysis of GWAS (MTAG)¹²⁸ to increase the precision of the estimates of our GWAS of self-reported general risk tolerance and to improve our ability to detect lead SNPs associated with general risk tolerance. MTAG offers several advantages over lookup-based methods, such as the proxy-phenotype method (**Supplementary Information section 8**); MTAG allows for more than two phenotypes, it increases detection power for all included phenotypes, and it works even in the presence of sample overlap across the various GWAS.

With MTAG, we leveraged the additional information contained in the GWAS summary statistics of six phenotypes related to risk tolerance or risky behavior. The six phenotypes are the five supplementary GWAS phenotypes adventurousness, automobile speeding propensity, drinks per week, ever smoker, and number of sexual partners, as well as lifetime cannabis use^{100,mm,nn}. We selected these six phenotypes because they each plausibly capture a different dimension of risk-taking behavior, and they all are significantly genetically correlated with general risk tolerance (**Supplementary Table 7.1, Panels A1 and A3**). Additionally, we conducted an additional MTAG analysis that also included self-employment¹⁰¹ among those phenotypes, but ultimately decided not to include self-employment in our baseline analysis, as we explain below.

We also used the summary statistics of this MTAG analysis to construct polygenic scores of general risk tolerance and evaluate their power in predicting a suite of phenotypes available in three validation cohorts (**Supplementary Information section 10**).

9.2 Methods

We used the summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance and of the six aforementioned phenotypes^{oo} as input for our MTAG analysis.

MTAG builds on the assumption that the correlation in the effect size of a SNP across phenotypes is the same for all SNPs. This assumption is strong and often violated; however, Turley *et al.* (2017)¹²⁸ analytically show that, as long as the SNPs have a non-null association with all phenotypes, MTAG is still a consistent estimator with a lower mean squared error than a corresponding GWAS. A problem might arise in MTAG for SNPs that have no effect on one phenotype but a sizeable effect on another. This might be the case for SNPs in the proximity of

^{mm} The summary statistics for most of these phenotypes come from GWAS that were conducted in samples that included the UKB and 23andMe cohorts. This should not bias the results because, as mentioned above, MTAG works well in the presence of sample overlap across the various GWAS.

ⁿⁿ We do not use the summary statistics from our sixth supplementary GWAS phenotype, the first PC of the risky behaviors, because that would have been redundant given that we already use the summary statistics of the four risky behaviors.

^{oo} As usual, we applied genomic control using the intercepts of LD Score regressions to adjust the summary statistics used as input for our MTAG analysis. As the MTAG summary statistics are invariant to any scaling performed on the summary statistics used as inputs for the analysis, whether genomic control has been applied to the input summary statistics should not affect the results. We did not apply genomic control using the intercepts of LD Score regressions to adjust the summary statistics outputted by the MTAG analysis, because the adjustment is already built into the MTAG estimates.

genes implicated in a biological process that is specific to one of the phenotypes, but unlikely to be implicated for general risk tolerance. Such examples might include nicotine or cannabis receptors, or alcohol metabolism. We therefore excluded from this analysis all SNPs within 1Mb of the genes *CHRNA5* and *CHRN3* (nicotinic receptors), *CNR1* and *CNR2* (cannabinoid receptors), and *ADH1B* (Alcohol Dehydrogenase)^{pp}.

We imposed a MAF filter of 0.01 and a sample size filter $N \geq \frac{2}{3} \times P_{90}(N)$ to the SNPs for all datasets, where $P_{90}(N)$ denotes the ninth decile of the sample size of a dataset. MTAG automatically limits the analysis to the 5,869,552 SNPs present in all data sets.

We also considered adding self-employment¹⁰¹ to the seven phenotypes in the MTAG analysis, but doing so would have limited the number of shared SNPs to 2,232,479. Below, we also report (in parentheses) the results of the MTAG analysis that also included self-employment (and that therefore included a total of eight phenotypes).

We ran MTAG (July 13, 2017 release) and then clumped the resulting MTAG summary statistics for the general-risk-tolerance phenotype using our main reference panel³⁹ and PLINK 1.9, with the same thresholds we used for the main GWAS analysis (**Supplementary Information section 2.8**). These thresholds included a primary P value threshold (5×10^{-8}), a secondary P value threshold (1×10^{-4}), an r^2 threshold (0.1), and a SNP window defined in kilobases (1,000,000 kb)^{qq}.

To assess whether the lead SNPs from the MTAG analysis of general risk tolerance are in loci that have not been identified by previous GWAS of risk tolerance, we repeated the steps we had followed to assess the novelty of the lead SNPs from our discovery GWAS of general risk tolerance (described in **Supplementary Information section 2.10**).

9.3 Results

9.3.1 MTAG summary statistics

Leveraging the information present in the seven (eight with self-employment) selected phenotypes, MTAG increases the number of independent SNPs reaching genome-wide significance for general risk tolerance from 124 to 312 (225 with self-employment). **Supplementary Table 9.1** reports the 312 lead SNPs. 127 of these are in weak LD (pairwise $r^2 > 0.1$) with the 124 lead SNPs already identified in our discovery GWAS of general risk tolerance. The top SNP is located in the region in the *CADM2* gene on chromosome 3 that has previously been identified by Day *et al.*⁶⁶ and that has also been associated in a concurrent study on general risk tolerance by Strawbridge *et al.*⁶⁷. A total of 185 of the 312 lead SNPs are novel genome-wide significant associations with general risk tolerance that have not been identified by our discovery GWAS, by Day *et al.*, or by Strawbridge *et al.*

^{pp} It is worth noting that SNPs close to the *ADH1B* gene were top-hits in recent GWAS of alcohol consumption (Clark *et al.* (2017)²⁷² and **Supplementary Table 3.2**). By contrast, SNPs in the proximity of the cannabis receptor genes were not associated with life-time cannabis use at the genome-wide significant level in the GWAS of lifetime cannabis use whose summary statistics we obtained¹⁰⁰. SNPs close to the nicotine receptors genes have been found to be significantly associated with daily cigarettes smoked but not with ever smoker (TAG *et al.* (2010)³⁵ and **Supplementary Table 3.2**), although this could be due to the limited statistical power of those GWAS. For precautionary reasons, we still drop those SNPs from the MTAG analysis. None of these regions contain top-hits for the single-trait GWAS of general risk tolerance, and therefore their exclusion should have little bearing on our analysis.

^{qq} As noted before, we used a very wide SNP window of 1,000,000 kb, which effectively makes the r^2 and P value thresholds the only binding parameters for the PLINK clumping algorithm.

MTAG increases the mean χ^2 for general risk tolerance from 1.81 to 2.21 (1.89 to 2.23 with self-employment). To achieve a similar increase in χ^2 , the sample size for the GWAS would have to be increased from 975,353 to 1,452,014 (975,353 to 1,346,482 with self-employment). This increase in signal can be seen by comparing the Q-Q plots for the MTAG analysis with the seven phenotypes and for the meta-analysis of the discovery and replication GWAS of general risk tolerance (**Extended Data Fig. 9.1a**).

Extended Data Fig. 9.1b displays the Manhattan plots for the MTAG analysis of general risk tolerance (top panel) and for the discovery GWAS of general risk tolerance (bottom panel), revealing strong similarities. Indeed, the genetic correlation between the summary statistics from the MTAG analysis and from the meta-analysis of the discovery and replication GWAS of general risk tolerance is very close to unity ($\hat{r}_g = 0.959$, $SE = 0.0025$ without self-employment; $\hat{r}_g = 0.975$, $SE = 0.0016$ with self-employment).

9.3.2 *Robustness tests*

MTAG can lead to inflated results if the summary statistics for some of the phenotypes come from highly-powered GWAS but have relatively low genetic correlations with the phenotype of interest. Here, this might be the case for the ever smoker phenotype, since the summary statistics of that phenotype have a mean χ^2 statistics of 2.006 but a genetic correlation of only 0.28 with general risk tolerance. However, we repeated the above MTAG analysis without the ever smoker phenotype (and without self-employment), and we still found 299 independent SNPs that reach genome-wide significance, 296 of which are either the exact same SNPs or within loci of the original MTAG analysis. The remaining three SNPs have P values close to our P value threshold of 5×10^{-8} .

9.3.3 *Predictive power of general-risk-tolerance polygenic score constructed with the MTAG summary statistics*

We constructed a polygenic score using the summary statistics from the MTAG analysis that used the seven selected phenotypes. This score allows us to measure how the increase in signal translates into increased out-of-sample predictive power for several measures of risk tolerance, personality traits, and risky behaviors in the Add Health and the HRS cohorts. The construction of the polygenic score with the LDpred method¹²⁹ (with the Gaussian mixture weight 0.3), the definition of the predicted phenotypes, and the results of these analyses are described in detail in **Supplementary Information section 10**. Comparison of the predictive power of the polygenic scores constructed using the MTAG and GWAS summary statistics are reported in **Extended Data Figs. 10.1-10.2** and in **Supplementary Tables 10.1-10.4**. As expected, the predictive power of the polygenic scores constructed using the MTAG summary statistics is on average slightly higher than that of the scores constructed using only the GWAS summary statistics.

10 Polygenic prediction

To assess the out-of-sample predictive power of the genetic variants associated with self-reported general risk tolerance, and to gauge the potential for leveraging these associations in empirical research in the behavioral sciences, we constructed various polygenic scores (PGS) and used them to predict several risk-related phenotypes, personality traits, and real-world measures of risky behaviors.

In this section, we first describe in detail the phenotypes we predicted, which fall in three main domains: measures of risk tolerance (including general risk tolerance), personality traits, and risky behaviors. To be able to analyze several phenotypes in all of these domains, we used six different datasets which contain rich phenotypic information: the Add Health, HRS, NTR, STR^{rr}, UKB-siblings^{ss}, and Zurich cohorts.

Next, we delineate and motivate the methodology that we followed to construct the polygenic scores used for prediction. We constructed three polygenic scores in total. Our first two polygenic scores were constructed with the LDpred¹²⁹ method, which accounts for the linkage disequilibrium (LD) between SNPs. The first was constructed using the summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance; the second was constructed using the MTAG summary statistics (for details, see **Supplementary Information section 9**). Our third polygenic score was constructed with the classical method, which simply weights SNPs by their GWAS effect size^{130,131}. Due to data access limitations, the 23andMe cohort could not be included in the meta-analysis whose summary statistics we used to construct the polygenic scores in the NTR, STR, and Zurich cohorts. The polygenic score using the MTAG summary statistics was only constructed for the Add Health, HRS, and UKB-siblings cohorts.

Across our analyses, we find that the polygenic scores' predictive power is within the range expected according to theory^{132,133}, when we take into account the SNP heritability of general risk tolerance and cross-cohort heterogeneity. In addition, our polygenic scores are predictive of both general risk tolerance and alternative measures of risk tolerance, such as financial and income gamble risk tolerance. Furthermore, the scores are predictive of a wide variety of other phenotypes: from personality traits such as openness to experience and behavioral inhibition, to real-world economic behaviors such as being an entrepreneur or owning a business, to risky health behaviors such as drinking, smoking, or drug use, to precautionary behaviors such as having life and health insurance. These results are robust to controlling for educational attainment, cognitive performance, and personality traits. Finally, for some of the predicted phenotypes, the predictive power of our polygenic scores is comparable to, or even exceeds, the predictive power of cognitive performance or educational attainment.

^{rr} The STR1 and STR2 cohorts were merged for the prediction analysis. See the Appendix at the end of this section for details.

^{ss} As mentioned in **Supplementary Information section 4**, the UKB-siblings cohort was defined in the same way as the full UKB cohort, but only includes individuals with at least one full sibling in the UKB.

10.1 Phenotype definition

10.1.1 Risk-tolerance phenotypes

We predict both general-risk-tolerance measures and several alternative measures of risk tolerance. These alternative measures have been developed to elicit risk tolerance in specific domains, such as financial investment decisions or career choices. Three such alternative measures are available in the cohorts we analyze: “financial risk tolerance,” “income gamble risk tolerance,” and “lottery-elicited risk tolerance.” These alternative measures have been shown to correlate with non-hypothetical risky behaviors such as smoking and drinking, being self-employed, or holding a risky investment portfolio^{1–5}. As a negative control, we also predicted height in the STR cohort.

(1) General risk tolerance

The general-risk-tolerance phenotypes we used are comparable to the measures used in the discovery and replication GWAS of general risk tolerance (these measures are described in **Supplementary Information section 1.2**).

In the STR cohort, the following question adapted from the German Socioeconomic Panel⁴ was asked to the respondents: *“How do you see yourself: Are you generally a person who is fully prepared to take risks or do you try to avoid taking risks? Please tick a box on the scale, where the value 1 means ‘unwilling to take risks’ and the value 10 means ‘fully prepared to take risks’.”* The sample size for general risk tolerance in the STR cohort is 8,012, and the mean response is 4.5 (**Supplementary Table 10.1**).

In the Add Health cohort, the latest wave asks respondents who were 24–34 years old the following question: *“How much do you agree with each statement about you as you generally are now, not as you wish to be in the future?: I like to take risks.”* Likert-scale response options include: [1] strongly agree; [2] agree; [3] neither agree nor disagree; [4] disagree; [5] strongly disagree. We reverse coded this variable, so that individuals more likely to take risks were coded with a “5” rather than a “1,” and vice versa. In total, we have data available for 4,749 European respondents for this variable, and the mean response is 3.0 (**Supplementary Table 10.1**).

In the UKB-siblings cohort we have information on self-reported general risk tolerance. All of the phenotypes predicted in the UKB-siblings cohort are the same as the UKB measures used in our primary and supplementary GWAS. For a detailed description of these phenotypes, see **Supplementary Information section 1.2**.

(2) Financial risk tolerance

Financial risk tolerance was measured with the following survey question in the STR cohort¹⁰:

“Are you a person that is fully prepared to take financial risk or do you try to avoid taking financial risk? Please tick a box on the scale, where the value 1 means: ‘not at all willing to take risks’ and the value 10 means: ‘very willing to take risks’.”

The sample size for financial risk tolerance in the STR cohort is 8,038, with a mean response of 3.5 (**Supplementary Table 10.1**).

(3) Income gamble risk tolerance

The income gamble risk tolerance phenotype is available in the STR and HRS cohorts. In the STR cohort the income gamble risk question is¹³⁴:

“Imagine the following hypothetical situation. You are the sole provider for your household, and you have the choice between two equally good jobs:

Job A will with certainty give you SEK 25,000 per month after taxes for the rest of your life.

Job B will give you a 50-50 chance of SEK 50,000 per month after taxes for the rest of your life, and a 50-50 chance of SEK X per month after taxes for the rest of your life.

Which job do you choose?”

The question was asked three times, and the amount X consecutively takes the values 20,000, 22,000 and 17,000. At the time, one US dollar was worth approximately seven SEK. A respondent receives one point for each time the risky Job B is chosen, so that a value of 0 indicates the lowest risk tolerance, and a value of 3 indicates the highest risk tolerance.

The sample size for income gamble risk tolerance in the STR cohort is 7,577, with a mean response of 2.0 (**Supplementary Table 10.1**).

In the HRS cohort, the income gamble risk question was asked in several waves. Here, we summarize how we created the income gamble risk tolerance variable in the HRS cohort. We provide further details in the Appendix at the end of this section.

In the first wave the income gamble risk question is phrased as follows:

“Suppose that you are the only income earner in the family, and you have a good job guaranteed to give you your current (family) income every year for life. You are given the opportunity to take a new and equally good job, with a 50-50 chance it will double your (family) income and a 50-50 chance that it will cut your (family) income by a third. Would you take the new job?”

If the answer is “yes” to the first question, then the respondent is asked this follow-up question:

“Suppose the chances were 50-50 that it would double your (family) income, and 50-50 that it would cut it in half. Would you still take the new job?”

If the answer is “no” to the first question, then the respondent is asked this follow-up question:

“Suppose the chances were 50-50 that it would double your (family) income and 50-50 that it would cut it by 20 percent. Would you then take the new job?”

In the following waves, the question wording is slightly modified, albeit keeping the same overall structure (see the Appendix at the end of this section for more details). In order to merge the information from different waves, we first coded each variable so that higher values imply higher risk tolerance, and then we took the average of the residuals from a separate OLS regression of each wave’s phenotype on an intercept and birth year, birth year squared, birth year cubed, sex, as well as three interaction terms between sex and the three birth year variables.

The sample size for income gamble risk tolerance is 7,302 in the HRS cohort, with a mean answer of 1.8 (**Supplementary Table 10.1**).

(4) Lottery-elicited risk tolerance

The lottery-elicited risk tolerance phenotype is only available in the Zurich cohort. It was elicited using a method called Multiple Price List (MPL)^{4,25,135}, which is commonly used in studies on risk tolerance based on incentivized gambles or lotteries. This phenotype was measured in an incentivized computerized experiment, in which respondents were asked lottery questions presented in three separate screens, corresponding to three tables. Each screen shows a table with 20 rows, on which respondents are required to make 20 binary decisions between a fixed lottery (option A) and a certain outcome (option B), which changes from row to row.

In each table, option A offers the possibility to “Win X CHF with a probability of 50% and win Y CHF with a probability of 50%.” The following values for X and Y are used for the three tables: Table 1: (20, 10) CHF; Table 2: (20, 0) CHF; and Table 3: (50, 20) CHF. (The expected values for option A are thus 15, 10, and 35 for the three tables). Option B displays the guaranteed win as the interval [X, Y] divided into 20 equal steps presented in descending order on each screen. (Between 10 and 19.5 CHF on table 1, between 0 and 20 CHF on table 2, and between 20 and 48.5 CHF on table 3.) Respondents chose between either option A or B for each of the 20 rows and the software enforced consistency so that respondents could only switch once from option B to option A on each screen. We calculated a screen’s certainty equivalent as the average of the two option B wins just above and below the respondents’ switching point. Individual i ’s lottery-elicited risk tolerance measure was then calculated as the median of the calculated relative risk premia (rrp) of the three lottery questions:

$$(8) \quad measure_i = median(rrp_{i,j}) = median\left(\frac{(B_{i,j} - E[A_j])}{|E[A_j]|}\right),$$

where $E[A_j]$ is the expected value of option A on screen j and $B_{i,j}$ is the certainty equivalent for individual i on screen j . (For the subsamples from Munich and Innsbruck, currencies were converted to EURO according to purchasing power.)

The sample size of the lottery-elicited risk tolerance is 2,610 in the Zurich cohort.

10.1.2 Personality traits

Personality traits have been defined as “the relatively enduring patterns of thoughts, feelings, and behaviors that reflect the tendency to respond in certain ways under certain circumstances”¹³⁶. Therefore general risk tolerance can be viewed as one of the many traits characterizing an individual’s personality, and is certainly very much related to established personality measures such as behavioral inhibition (the tendency to have a restrained or fearful response to unfamiliar events), openness to experience (the degree of intellectual curiosity, creativity, and preference for novelty and variety), or sensation-seeking (the tendency to search for experiences and feelings that are varied, novel, complex, and intense). Indeed several studies at the intersection of economics and personality psychology document the phenotypic correlation between measures of personality traits and general risk tolerance^{2-4,137-142}. Motivated by these findings, we investigate the predictive power of our genetic measure of risk tolerance for various phenotypic measures of personality traits.

A large range of measurements of personality traits is available in our datasets, from the widely used five-factor model of personality (the Big Five), to more specific measures like behavioral inhibition and locus of control. Below we provide a detailed description of our personality phenotypes, and we present the summary statistics in **Supplementary Table 10.2**.

(1) Behavioral inhibition

Behavioral inhibition is defined as a behavioral style characterized by a restrained or fearful response to unfamiliar events, both social and non-social, and has been shown to be moderately stable over time¹⁴³. The 16-item Adult Measure of Behavioral Inhibition (AMBI) battery is available in the STR cohort ($n = 7,608$) cohort. An example question is: “Do you feel awkward when you are approached by someone new?” Each question is answered on a three-point scale. Summing the responses leads to a variable taking values in the interval $[0, 32]$. We reverse coded the variable so that a higher score implies a higher behavioral *disinhibition*, since we expect behavioral disinhibition to be positively correlated with general risk tolerance, as has been found in previous studies².

(2) Locus of control

Locus of control is a personality trait defined on a spectrum ranging from internal control, where an individual feels that the outcomes of events are contingent on his or her own behavior and attributes, to external control, where an individual believes that external forces outside of personal control determines the outcome of events. The 12-item locus of control scale is available in the STR cohort for 6,777 individuals. It is coded such that higher numbers indicate an internal locus of control. An example question is: “Becoming a success is a matter of hard work; luck has little or nothing to do with it”¹⁴⁴. We expect internal locus of control to be positively correlated with general risk tolerance, and this is what has been found in previous studies².

(3) Big Five personality traits

Several decades of research in personality psychology beginning in the 1970s have led to a widely shared taxonomy of personality traits, known as the Big Five^{145–147}. Identified via factor analysis of common language descriptors of personality, these five distinct and independent personality characteristics are openness to experience, conscientiousness, extraversion, agreeableness, and neuroticism. Such traits have been measured both in the HRS and in the NTR cohorts.

In the HRS cohort, the Big Five personality traits were measured in four waves between 2006 and 2012, with 26 items in 2006–2008 and 31 in 2010–2012. The original 26 items in 2006 and 2008 were obtained from the MIDUS survey¹⁴⁸. Extraversion, agreeableness, and conscientiousness were measured with five items, openness to experience with seven, and neuroticism with four. In 2010 and 2012, five items from the International Personality Item Pool¹⁴⁹ were added for conscientiousness. As is common in the literature, the responses to each item were elicited on a four-point scale¹⁴⁶. For each trait, scores were constructed by taking the mean of all items in the respective category after recoding items to make their direction consistent, and then averaged across waves and standardized to have zero mean and unit variance. A score was set to missing if more than half of the items in that category had missing values. The resulting sample sizes are 8,253 for openness, 8,268 for conscientiousness, 8,271 for extraversion, 8,271 for agreeableness, and 8,264 for neuroticism. Given the limited evidence on the sign of the phenotypic correlation

between general risk tolerance and the Big Five traits, we do not have an *ex ante* expectation about the signs of their associations with our polygenic score of general risk tolerance.

In the NTR cohort, the NEO Five-Factor Inventory (NEO-FFI) was sent to adult participants in three waves in 2003, 2009 and 2013. Each of the five personality traits is measured with 12 items, and the responses to each of these were given on a five-point Likert scale. Scores were standardized to have zero mean and unit variance in each wave and then averaged across waves, yielding a sample size of 8,526.

(4) Sensation seeking

Sensation seeking is defined by the search for experiences and feelings that are “varied, novel, complex and intense,” and by the readiness to “take physical, social, legal, and financial risks for the sake of such experiences”¹⁵⁰.

In the NTR cohort, sensation seeking was assessed using a short version of the Zuckerman sensation seeking scale (SSS)^{151–153}. The scale consists of 12 items that measure four subdomains of sensation seeking. Thrill and adventure seeking ($n = 8,649$), defined as the desire to engage in extreme sports and risky activities; experience seeking ($n = 8,185$), defined as the desire for experiences that engage the mind and senses such as art, travel, food or dress; disinhibition ($n = 8,618$), defined as the desire to find release through interactions with other disinhibited people, wild parties and sexual disinhibition; and boredom susceptibility ($n = 8,621$), defined as the intolerance to boredom. A combined total score is then constructed as the total sum of all subscales. The SSS was measured at six time points between 1991 and 2007. Scores were standardized to have zero mean and unit variance in each wave and then averaged across waves.

(5) Attention-deficit hyperactivity disorder

We also examined the predictive power of our score for attention-deficit hyperactivity disorder (ADHD). ADHD is not a personality trait, but it is nonetheless of interest because it has been shown to be phenotypically correlated with risk-taking behaviors, such as starting one’s own business^{154–156}. ADHD is a chronic condition of persistent behavioral problems that often begins in childhood and persists into adulthood. The fourth edition of the Diagnostic and Statistical Manual of Mental Disorders (DSM-IV) defines ADHD as “a persistent pattern of inattention and/or hyperactivity-impulsivity that is more frequently displayed and more severe than is typically observed in individuals at a comparable level of development.”

In the NTR cohort, symptoms of ADHD were assessed by the Conners ADHD index^{157,158} and were measured at three occasions in 2004, 2007 and 2013. The scores were standardized to have zero mean and unit variance in each wave and averaged across waves, yielding a total of 8,457 respondents.

10.1.3 Risky behaviors

Building on previous studies that demonstrate the predictive power of risk tolerance for various behaviors^{1–5,28}, we consider a wide range of phenotypes associated with lifestyle risks. We are interested in two categories of real-world behaviors: *risky* and *precautionary*. The first category focuses on behaviors that increase an individual’s exposure to unpredictable outcomes and includes smoking, drinking, unhealthy food consumption, financial investment, and career

choices. The second category focuses on behaviors that aim to prevent or curtail potential future negative outcomes, and includes investing in health or life insurance as well as preventative health care measures such as taking medications and getting regular medical screenings.

Below we provide a detailed description of our real-world phenotypes. **Supplementary Table 10.3** presents summary statistics. For a more detailed description of measures drawn from the STR cohort, see Beauchamp *et al.* (2015)².

(1) Smoking

Smoking is associated with increased risks of heart disease, stroke, lung cancer (and other types of cancers), and other chronic lung diseases¹⁵⁹. Smoking has also been shown to be phenotypically correlated with risk tolerance¹⁻⁵.

In the HRS cohort, we constructed a binary variable for “ever smoker” that equals “1” if an individual respondent reported ever having been a tobacco smoker in at least one wave. 57% of the 8,617 HRS respondents are coded as ever smokers.

In the STR cohort, respondents are coded as “ever smokers” if they indicated they currently smoke regularly, used to smoke regularly, currently smoke on and off, or used to smoke on and off. 53% of the 13,921 respondents in the STR cohort are coded as smokers.

In the Add Health cohort, respondents are coded as “ever smokers” if they reported having ever smoked regularly (i.e., at least one cigarette every day for 30 days) in Wave 4, the latest wave of the Add Health data. 52.6% of 4,775 European ancestry respondents were coded as smokers. We also constructed a continuous variable for age of smoking initiation among those who were coded as smokers, again in Wave 4. The mean age of smoking initiation in the Add Health cohort is 16.9, and the variable is defined for 2,493 European respondents.

In the UKB-siblings cohort we have information on whether the respondent has ever been a smoker. For a detailed description of the phenotype used, see **Supplementary Information section 1.2**.

(2) Drinking

Similar to smoking, drinking is a risky behavior known to have negative health consequences¹⁶⁰ and has also been correlated phenotypically with risk tolerance^{1,2}. The Add Health, HRS, and STR datasets include a rich set of questions on drinking habits.

In the HRS cohort, we calculated the number of alcoholic drinks per week reported by each respondent in each wave, took the average across waves, and normalized the resulting measure in a variable with a mean of approximately zero and a variance of approximately one^{tt}. Importantly, in waves 1 and 2, respondents were only able to choose among 5 categories: 0 drinks per week, 1 or fewer drinks per week, 1-2 drinks per week, 3-4 drinks per week, or 5 or more drinks per week. We coded these as 0, 1, 1.5, 3.5, and 5 drinks per week for each wave, respectively. From wave 3 onward, respondents were first asked how many days per week they had *any* alcoholic drinks, and

^{tt} The normalized variable is the inverse normal cumulative distribution function (CDF) of the respondents’ percentile ranks. Given the discrete nature of the underlying drinking reports, the inverse normal CDF has some mass points and does not have a mean of exactly zero and a variance of exactly one. This is true of all the other phenotypes that have been normalized.

then were asked how many drinks they had on average on each of those days. For these waves, we simply multiplied each of these two responses to calculate drinks per week. This information was available for 8,652 respondents.

Similarly, in Add Health we calculated the number of drinks per week for 3,712 European respondents in Wave 4. Respondents were asked both how many days they consumed alcohol in the last 12 months *and* how many alcoholic drinks they consumed each time they drank in the last 12 months. We multiplied these responses together and divided by 52 to arrive at our final drinks per week variable. The average number of drinks per week in Add Health was 5.6.

In the UKB-siblings cohort we have information on the number of drinks per week. For a detailed description of the phenotype used see **Supplementary Information section 1.2**. Notice that drinks per week was normalized in the HRS but not in the Add Health and UKB-siblings cohorts, where it is simply reported as a count.

In the STR cohort, we constructed a binary variable equal to “1” if respondents reported having alcoholic drinks more than twice in the past month. Specifically, a first question asks respondents if they drank strong beer, wine, or liquor more than twice in the past month. If respondents answer in the negative, a follow-up question asks about their frequency of drinking for each type alcohol separately, and “twice per month” was one of eight response categories. Overall, we classified individuals as alcohol consumers if they answered the first question in the affirmative or if their answers to the follow-up questions indicated that they usually drink beer, wine, or liquor at least twice a month. 82% of the 12,551 respondents in the STR cohort were coded as such.

The STR dataset contains additional information regarding excessive drinking over the life course, which might better capture risky drinking behavior. We constructed a dummy variable equal to “1” for respondents who reported any of the following: 1) ever having a period in their life when they drank too much; 2) ever having a period in their life when someone else objected to their drinking; or 3) ever having a period in their life when they chose drinking instead of working, spending time on hobbies, or spending time with family or friends. 8% of the 13,269 STR respondents were coded as excessive drinkers.

(3) Cannabis consumption

Similar to cigarettes and alcohol, cannabis abuse can lead to negative health¹⁶¹ and social consequences¹⁶², especially during adolescence and early adulthood. Furthermore in 2008, at the time of reporting, recreational use of marijuana was illegal in the United States, and therefore its consumption carried the additional risk of breaking the law.^{uu}

Using data from Wave 4 of Add Health, we constructed a binary variable for “ever cannabis” that equals “1” if an individual respondent reported having ever used cannabis. 60.0% of the 4,742 European respondents had a value of “1” for this variable.

^{uu} Starting with Oregon in 1973, the possession of marijuana was decriminalized by some states, but remained illegal across the United States. Starting with California in 1996, some states legalized the use of marijuana for medical purposes. Recreational use was legalized only starting with Colorado in 2012. While the negative health and social consequences were still prevalent through the sample, the legal consequences of cannabis consumption varied depending on the residence of the respondents, which is unknown to us.

(4) Unhealthy food consumption

Consumption of fried food is a common risky health behavior that has been linked to cardiovascular disease and other negative health consequences¹⁶³. Questions about fried food consumption were asked to a subset of 3,078 respondents in the HRS cohort who participated in the 2013 Healthcare and Nutrition Mailout. We looked at propensity to eat fried food either at home or as takeout. In reporting how much fried food they consumed, respondents could answer “Never,” “Less than once per week,” “1-3 times a week,” “4-6 times a week,” or “Daily.” We normalized the measures separately for home and for takeout, resulting in two measures with means of approximately zero and variances of approximately one. Our measure of overall propensity to eat fried food is the sum of these two normalized measures.

(5) Number of sexual partners

As a final measure of risky health behavior, we consider the total number of sexual partners, which is a risk factor for contracting sexually transmitted infections (STIs)^{164,165}. In Wave 4 of Add Health, respondents were asked with how many female and male partners they had ever engaged in any type of sexual activity. We summed the values for female and male partners. The mean number of sexual partners for the 4,603 European respondents was 13.5.

In the UKB-siblings cohort we have information on the number of sexual partners. For a detailed description of the phenotype used, see **Supplementary Information section 1.2**. Notice that number of sexual partners was normalized in the UKB-siblings cohort but not in the Add Health cohort, where it is simply reported as a count.

(6) Automobile speeding propensity

In the UKB-siblings cohort we have information on automobile speeding propensity. For a detailed description of the phenotype used, see **Supplementary Information section 1.2**.

(7) First PC of risky behaviors

Following the approach laid out in **Supplementary Information section 1.2**, we computed the first principal component (PC) of our four supplementary risky behaviors (automobile speeding propensity, drinks per week, ever smoker, and number of sexual partners) in the full UKB sample, and then extracted the individuals comprising the UKB-siblings cohort.

(8) Financial market participation

We consider two measures of exposure to financial risk, both of which have been correlated to phenotypic measures of risk tolerance^{1,2,4,28}. The first is financial market participation. This phenotype captures what economists refer to as the “extensive margin” of financial risk-taking behavior (i.e., does one participate in the financial market or not).

In the HRS cohort, we coded a dummy variable that equals “1” if a respondent ever reports a strictly positive “net value of stocks, mutual funds, and investment trusts” across all available waves. Using this measure, 68% of 8,652 HRS respondents participated in financial markets.

Similarly, in the STR cohort we constructed a variable coded “1” if respondents reported a positive value of either stock or bond holdings, using information from the single wave in this cohort when assets were reported. Overall, 30% of the 3,285 STR respondents were coded as participating in the financial market.

(9) Equity share and share of financial assets

Our second measure of exposure to financial risk considers the risk profile of respondents’ total wealth. We use the share of respondents’ total wealth held in equity (i.e., stocks) and similar risky financial assets. This phenotype captures what economists refer to as the “intensive margin” of financial risk-taking behavior (i.e., how much of a person’s wealth is exposed to financial market fluctuations).

In the HRS cohort, we calculated for each wave the ratio of “net value of stocks, mutual funds, and investment trusts” over “total wealth including secondary residence.” For each respondent, we then averaged this measure across all available waves. This measure roughly reflects the amount of total wealth an individual holds in risky financial assets. We note that mutual funds and investment trusts might include equities, as well as other risky assets such as bonds and derivatives. We refer to this resulting variable as the “share of financial assets.” In total, we dropped 50 observations reporting negative equity holdings and two observations for which the share of financial asset in a wave exceeds unity. The mean value of the share of financial assets variable across the 8,599 respondents is approximately 8%.

In the STR cohort, we instead calculate equity share as the ratio of the value of assets held in stocks to the sum of the value of assets held in property, stocks, bonds, bank, boat, and other assets such as jewelry, antiques, and art. These data are only available for a subset of the respondents because the associated questions were removed from the survey in the later waves (in an effort to reduce survey length). The 3,170 respondents in STR cohort held on average 3.5% of their total wealth in stocks.

(10) Career choices

While some careers are relatively secure and provide a steady stream of income, others are riskier and have greater income fluctuations. Prominent examples of the latter are owning a business or becoming an entrepreneur.

In order to capture this domain of risk taking, which has been correlated to phenotypic measures of risk preferences^{2,4,5,28}, we considered two separate measures in the STR cohort. First, we constructed a binary variable equal to “1” if respondents reported ever running their own business. 25% of the 7,979 individuals in STR cohort were coded as having ever run their own business.

As a second measure, we consider a self-report of being an entrepreneur, defined as someone who “commercializes a new innovation or idea ... [and who] has, or plans to have, a number of employees and strives to expand the business,” as opposed to a “self-employed person [who] owns and runs his/her own company, for instance a restaurant or a law firm, where he/she works [and] ... normally does not strive to expand over a certain limit and has 0 or a few employees.” This question was asked only of respondents who had reported ever running their own business. We constructed a binary variable equal to “1” if respondents reported being entrepreneurs, and “0” if the respondents reported being self-employed persons or if they reported never having run their

own businesses. Based on this measure, 5% of the 7,981 individuals in STR cohort were coded as entrepreneurs.

(11) *Preventative healthcare and healthcare utilization*

Focusing now on precautionary behaviors, we first consider choices regarding preventive healthcare.

We constructed an index of preventative health care and health care utilization by combining several available measures for 8,648 respondents in the HRS cohort. Specifically, we considered whether respondents had visited a doctor in the last two years, whether respondents had visited a dentist in the last two years, and whether respondents had recently undergone a number of specific procedures or screenings. These included cholesterol screening, flu shot, breast exam (for females), mammogram (for females), pap smear (for females), and prostate exam (for males). We took the simple sum of dummy indicators for having undergone or participated in any of these eight healthcare measures. Since not all of these items apply to both sexes, we normalized these sums separately for men and women, and then combined the measures, obtaining an index with a mean of approximately zero and a variance of approximately one.

(12) *Health insurance*

In order to limit the risk associated with an unexpected negative health event, individuals can enroll in health insurance. Prior to the implementation of The Patient Protection and Affordable Care Act (popularly known as “Obamacare” or “Healthcare reform”) in the US, this form of risk prevention was an individual choice (rather than required by law) for most individuals. In the HRS cohort, we measure whether individuals choose to have health insurance. As is the case for our other phenotypes, risk tolerance has been shown to predict whether a person has health insurance¹.

According to national and state eligibility criteria, some HRS respondents in certain waves were automatically granted health insurance (i.e., Medicaid or Medicare recipients and CHAMPUS/VA healthcare recipients), and therefore these observations are not used in the construction of this measure for these respondents. For the remaining observations, we calculated whether each respondent-wave observation responded affirmatively to questions about having each of several forms of health insurance (e.g., non-Medicare and non-CHAMPUS/VA federal health insurance, health insurance from current or previous employers, health insurance from spouse’s current or previous employers, other health insurance). For each respondent, we then took the average of non-missing observations across all waves until (and including) the 2011 wave^{vv}. Our resulting measure indicates the percentage of waves during which a respondent had healthcare, among waves during which it was an optional choice. On average, the 8,454 respondents in the HRS cohort were covered by health insurance 95% of the time.

^{vv} The individual mandate of the Affordable Care Act of 2010 was not yet in effect during 2011, the final wave from which we draw these data.

(13) *Life insurance*

Finally, individuals purchase life insurance to mitigate the negative consequence of their deaths for their families and loved ones. Phenotypic risk tolerance has been shown to predict whether a person has life insurance¹.

We measured whether respondents are consistently covered by life insurance by calculating the percentage of waves during which HRS respondents reported having life insurance. On average, the 8,652 respondents in the HRS cohort report being covered by life insurance more than two-thirds (71%) of the time.

10.2 Methods

10.2.1 *Polygenic scores*

We constructed three polygenic scores. We constructed our first polygenic score with the LDpred¹²⁹ method, using summary statistics from our GWAS of general risk tolerance; we also constructed our second score with the LDpred method, but using the summary statistics of general risk tolerance outputted by the MTAG analysis (**Supplementary Information section 9**); and we constructed our third score with the classical method, using summary statistics from our GWAS of general risk tolerance. We will refer to these scores as LDpred-GWAS, LDpred-MTAG, and Classical-GWAS scores respectively.

Due to data access limitations and to avoid overfitting, different cohorts were included in the GWAS whose summary statistics were used to construct the scores. In the Add Health and HRS cohorts, the summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance were used to construct the scores, with no cohort excluded ($n = 975,353$). For the analyses in the NTR and Zurich cohorts, the scores were constructed with the summary statistics from a meta-analysis that excluded the 23andMe cohort (due to data access limitations, the sample size of the resulting meta-analysis was $n = 466,571$). For the analysis in the STR cohort, we excluded the 23andMe cohort (due to data access limitations) as well as the STR cohort itself (to avoid overfitting⁹⁵) from the meta-analysis whose summary statistics we used to construct the scores (the sample size of the resulting meta-analysis is $n = 458,558$)^{ww}. And for the analysis in the UKB-siblings cohort, to avoid overfitting⁹⁵ we reran our discovery GWAS after excluding all individuals in the UKB-siblings cohort from the UKB cohort (the sample size of the resulting meta-analysis was $n = 937,353$)^{xx}.

The prediction analyses with our LDpred-MTAG score was only conducted in the Add Health, HRS, and UKB-siblings cohorts, since these were the only three cohorts for which 23andMe data could be used, due to data access limitations.

Both the LDpred and the classical polygenic scores are calculated as the weighted sum of the M individual genotypes:

^{ww} We also note that, since the HRS, NTR, and Zurich cohorts do not contain data on general risk tolerance, none of these cohorts was included in any meta-analysis of general risk tolerance.

^{xx} No single individual is present in both the UKB-siblings cohort and in the “UKB-nonsibs” cohort (which we define as the subset of individuals we used for the discovery GWAS we reran for the prediction analysis in the UKB-siblings cohort). However, some individuals in the UKB-siblings cohort have relatives in the UKB-nonsibs cohort, and that this could lead to overfitting⁹⁵. We further discuss this in **Supplementary Note section 10.3.5**.

$$(9) \quad \hat{S}_i = \sum_{j=1}^M \hat{\beta}_j g_{ij} ,$$

where \hat{S}_i denotes the polygenic score of individual i , $\hat{\beta}_j$ is the estimated additive effect size of the effect-coded allele at SNP j , and g_{ij} is the genotype of individual i at SNP j (coded as having 0, 1, or 2 instances of the effect-coded allele)¹³⁰.

For the LDpred scores, the estimated additive effect sizes (the $\hat{\beta}_j$'s) are the estimates from the summary statistics adjusted for LD between the SNPs. To calculate LDpred scores, an LD reference file and a validation reference file must be provided. For HRS, STR, UKB-siblings, and the Zurich cohorts we used the 1000 Genomes-imputed data (Phase 1, Version 3) of the HRS^{yy} cohort as the reference sample. For the Add Health cohort, we used the HRC (Haplotype Reference Consortium) Genomes-imputed data (Version 1.1) of Add Health (n=4,775) as the reference sample. For the NTR cohort, the reference sample used for the construction of the LDpred scores consists of all five European populations from the 1000 Genomes dataset: Utah Residents (CEPH) with Northern and Western European Ancestry, Finnish, British, Iberian, and Toscani individuals ($n = 381$).

The LDpred method relies on a Gaussian mixture weight that corresponds to the assumed fraction of SNPs that are causal. We used the software called LDpred¹²⁹ to generate a score for each of the following mixture weights: 1, 0.3, 0.1, 0.03, 0.01, 0.003, 0.001, 0.0003, and 0.0001, using the individuals' best guess data and using only SNPs in the HapMap consortium phase 3 release^{166,167} with a MAF > 0.01 and an imputation quality of more than 0.7. We report only the analyses with the LDpred scores constructed with the Gaussian mixture weight 0.3, as these scores tended to perform better both across cohorts and predicted phenotypes. Incidentally, a fraction of causal SNPs of 0.3 also happens to match closely our estimate of the fraction of non-null SNPs (π) of 0.29 in our estimation of the posterior distribution of the SNPs' true effect sizes in **Supplementary Information section 5.1.3**.

For the classical polygenic scores, the estimated additive effect size $\hat{\beta}_j$ for SNP j is the GWAS estimate for SNP j . We used the software PLINK⁴³ to produce the classical scores, using the individuals' best guess data; as with the LDpred method, we only use SNPs in the HapMap consortium phase 3 release¹⁶⁶ with a MAF > 0.01 and an imputation quality of more than 0.7 (the prediction results are thus comparable across the two methods)^{zz}.

^{yy} The HRS genotype data used as a reference sample for the STR and the Zurich cohorts was restricted according to the following quality control criteria. We removed 13,973 SNPs that have been flagged as having incorrect annotations from the HRS cohort²⁷³; we restricted the reference file to SNPs with imputation quality greater than 0.7, MAF greater than 0.01, SNP call rate greater than 95%; and we removed individuals with a call rate less than 95%, as well as related individuals and individuals not of North-Western European ancestry. The resulting sample contained 7,302 individuals.

For the HRS and UKB-siblings cohort, we restricted the reference file to HapMap3 SNPs with MAF greater than 0.01 and SNP call rates greater than 98%; we also removed individuals with a call rate less than 98%, related individuals, and individuals not of European ancestry. The resulting sample contained 8,353 individuals.

^{zz} The classical score is often calculated with an LD-pruning and P value thresholding procedure. The LD-pruning is meant to achieve independence between the predictive set of SNPs, and the P value threshold excludes SNPs that are not estimated to be significantly associated with the phenotype. This procedure often discards information that could

We expect our LDpred-GWAS score to have greater predictive power than our Classical-GWAS score, since the LDpred method takes into account the non-independence between SNPs. Further, since the MTAG summary statistics leverage the additional information contained in the GWAS of genetically correlated phenotypes, we expect our LDpred-MTAG score to have the greatest predictive power.

10.2.2 *Main prediction exercise*

We ran two separate Ordinary Least Squares (OLS) regressions for each phenotype, cohort, and score. The first regression includes only our baseline controls: sex, birth year, birth-year squared, birth-year cubed, as well as the interactions between sex and the three birth-year variables, and the first ten principal components (PCs) of the cohort-specific genetic relatedness matrix; in the NTR and STR cohorts, dummy variables that indicate the different genotype platforms were also included. The second regression is identical to the first one, except that it also includes the polygenic score. Since the scores have mean zero and unit variance, the estimated coefficients on the score represent the change in real-world outcomes associated with a one-standard-deviation increase in the polygenic score of general risk tolerance. Our measures of interest are the regression coefficients on the scores and the incremental R^2 (or pseudo- R^2) of the scores, defined as the difference between the R^2 (or pseudo- R^2) of the two regressions. The 95% confidence intervals for the incremental R^2 estimates are calculated with the bootstrap percentile method, with 1,000 bootstrap samples^{168,169}.

10.2.3 *Robustness to inclusion of additional control variables*

Are these polygenic scores of general risk tolerance potentially useful for the empirical researcher in the behavioral sciences? Since many social-science datasets contain rich information about the individual characteristics of the participants, it is important to understand whether the polygenic scores for general risk tolerance are still predictive of risky behaviors even after controlling for relevant variables that capture important individual characteristics. For this reason, when analyzing real-world risky behaviors, we also estimate the incremental R^2 (or pseudo- R^2) of the scores while controlling for cognitive performance, personality traits, and educational attainment in the baseline regression.

These traits were selected as additional controls because they could plausibly be important determinants of risky and precautionary behaviors, and because many of these traits are phenotypically correlated with risk tolerance². This robustness exercise also makes it easier to compare our prediction results with the existing literature, which usually controls for some of these personal characteristics^{6,135,170,171}.

Furthermore, we also calculated the incremental R^2 for cognitive performance, for personality traits, and for educational attainment, defined as the difference between the R^2 of a regression including only our baseline controls, and another regression including the baseline control and the traits of interest. These three incremental R^2 values provide the applied researcher with interesting benchmark comparisons, and enable us to understand whether the predictive power of our polygenic score is comparable to the predictive power of other important individual characteristics.

increase the predictive power if it were properly accounted for¹²⁹. Here, we do not use an LD-pruning and P value thresholding procedure to drop SNPs prior to constructing the scores.

For the Add Health cohort, we control only for educational attainment and a measure of verbal cognition, since information on personality traits is available only for a small subsample of individuals. The measure of verbal cognition is a modified version of the Peabody Picture Vocabulary Test which was collected in the first wave of Add Health, when participants were 12 to 20 years old. In this test, respondents have to select the illustration that best fits the meaning of the word that an interviewer has just read aloud. This computer-adapted test consisted of eighty-seven items, and scores were age-standardized.

For the HRS cohort, the cognitive performance measure is the average of measures from waves 2-10. In each of these waves, the cognitive performance measure is the sum of a total word recall summary score (based on immediate and delayed word recall tasks) and a mental status summary score (based on counting, naming, and vocabulary tasks)^{aaa}. The Big Five factors (defined above) were used as measures of personality traits^{bbb}, and years of education as a measure of educational attainment.

For the STR cohort, the standardized score on a military conscription test was used as a measure of cognitive performance, behavioral inhibition and locus of control were used as measures of personality traits, and years of education was used as a measure of educational attainment. The cognitive performance measure was merged using conscription data provided by the Military Archives of Sweden. Men were required by law to participate in military conscription around the age of 18. We use the stanine scores of four subtests of logical, verbal, spatial, and technical ability. Following Rietveld *et al.* (2014)¹⁰², we use the first principal component of these four stanine scores as the measure of cognitive performance. No measure of cognitive performance is available for females.

Finally, for the UKB-siblings cohort we control for educational attainment, neuroticism, and cognition. Our measure of neuroticism follows Okbay *et al.* (2016)¹⁸: it is constructed as a respondent's score on a 12-item version of the Eysenck Personality Inventory Neuroticism scale. Individuals must answer at least 10 out of 12 binary-response questions to be included, and questions that remain missing will be coded as 0. To obtain a measure of cognition, we used a test designed to measure fluid intelligence. The test consists of thirteen logic and reasoning questions and has a two-minute time limit. Each respondent took the test up to four times. We used the mean of the standardized score across the occasions on which the respondent took the test as our measure of cognition.

10.2.4 Methodology for binary and censored phenotypes

For the analysis of some real-world risky behaviors, we used specific statistical methods to model more accurately the distribution of the dependent variable. For binary outcomes, such as engaging in smoking, drinking, or participating in financial markets, we estimated Probit models. Instead of the estimated coefficient, which has no clear interpretation in a Probit model, we report the average marginal effect of the polygenic score. We report this as $\frac{\partial \Pr(y=1|PGS,X)}{\partial PGS} = \hat{\beta}_{PGS} \cdot \hat{E}[\phi(PGS \cdot \hat{\beta}_{PGS} + X\hat{\beta})]$, where $\hat{\beta}_{PGS}$ is the estimated coefficient on the polygenic score, ϕ is the Gaussian probability density function, $\hat{E}[\cdot]$ takes the sample average, PGS is the polygenic score, and X contains the control variables. In the absence of a meaningful R^2 measure for the Probit model, we

^{aaa} In Wave 2, these measures are present only for a subset of respondents (the AHEAD cohort).

^{bbb} See above for a detailed description of the personality variables in the HRS cohort.

used the McFadden pseudo- R^2 , $\left(\frac{l_0 - l_M}{l_0}\right)$, where l_0 is the log-likelihood of a model with only a constant, and l_M is the log-likelihood of the full model. Our measure of the incremental pseudo- R^2 is thus $\frac{l_X - l_{X,PGS}}{l_0}$, which indicates the difference between the log-likelihood of the model controlling for baseline controls and the score ($l_{X,PGS}$) and the log-likelihood of the model controlling only for the baseline controls (l_X), scaled by the log-likelihood of a model with only a constant (l_0). For equity share and share of financial assets phenotypes, we estimate tobit models instead of OLS regressions, to account for the point mass at zero (due to many respondents in our dataset not holding any stocks)^{ccc}. The same measure of incremental pseudo- R^2 , $\left(\frac{l_X - l_{X,PGS}}{l_0}\right)$, is reported^{ddd}.

The 95% confidence intervals for all of these estimates are calculated with the bootstrap percentile method, with 1,000 bootstrap samples^{168,169}.

10.3 Results

Results are presented in **Extended Data Figs. 10.1-10.2** and **Supplementary Tables 10.1-10.4**. For each predicted phenotype, cohort, and score, these tables report summary statistics for the regression sample, the estimated incremental R^2 of the score, and the sign and P value of the estimated regression coefficient on the polygenic score.

Overall, we find that our preferred polygenic score explains 1.01% to 1.78% of the variation in general risk tolerance, up to 1.4% of the variation in several personality traits, and up to 1.94% of variation in real-world behaviors. As we discuss in **Supplementary Information section 10.4**, our incremental R^2 estimates from the prediction of general risk tolerance fall within the range we expect based on theory^{132,133}.

Below, we focus our discussion on the results for the LDpred-GWAS score. As mentioned above, the results displayed for the LDpred scores are based on a Gaussian mixture weight of 0.3. We chose this weight because the corresponding scores consistently performed well across cohorts and phenotypes in our analyses. The results for the LDpred-MTAG and Classical-GWAS scores are generally similar, although the predictive power of the LDpred-MTAG score tends to be slightly higher and that of the Classical-GWAS score tends to be slightly lower.

10.3.1 Risk tolerance

In the UKB, the incremental R^2 of the LDpred-GWAS score is 1.62% (CI: 1.37% - 1.90%). In the Add Health cohort we estimate a much smaller predictive power, with incremental R^2 of 1.01%

^{ccc} The tobit model is a maximum likelihood estimator proposed by James Tobin²⁷⁴ that assumes a linear relationship between regressors X , a normally distributed error term ε , and a continuous latent variable y^* . The observed dependent variable is $y = y^*$ whenever $y^* > 0$ but is $y = 0$ otherwise. This model is used to account for censoring and a point mass at zero, as is observed for our measures of equity share and of share of financial assets.

^{ddd} Because the tobit model has a continuous likelihood, the McFadden pseudo- R^2 can sometimes be smaller than zero or greater than 1, which is nonsensical for an R^2 measure. Whenever that happened, such as in the case of share of financial assets in the HRS cohort when cognitive performance or educational attainment are controlled for, we reported “N.A.” in the table of results.

(CI: 0.57% - 1.62%). In the STR^{ccc} cohort we estimate an incremental R^2 of 1.05% (CI: 0.64% - 1.50%). Using the summary statistics from the MTAG analysis improves our predictive power: the incremental R^2 of the LDpred-MTAG score is 1.78% in the UKB-siblings cohort and 1.07% in the Add Health cohort.

The estimated regression coefficients for all scores have the expected sign and are all highly statistically significant (with P values < 0.0005), indicating that the scores significantly predict general risk tolerance out of sample.

(1) Alternative measures of risk tolerance

The predictive power of our polygenic score for alternative measures of risk tolerance is also significant, but limited in magnitude.

For the financial risk tolerance phenotype, the incremental R^2 of the LDpred-GWAS score in the STR cohort is 0.44%. For the income gamble risk tolerance phenotype, the incremental R^2 is 0.24% in the HRS cohort, and 0.34% in the STR cohort. When predicting either financial risk tolerance or income gamble risk tolerance, the coefficients on the LDpred-GWAS scores are always highly statistically significant.

By contrast, we find that our polygenic scores of general risk tolerance do not predict the lottery-based measure of risk tolerance in the Zurich cohort. For this measure, the incremental R^2 is 0.09% and the coefficient on the score is not distinguishable from zero ($P = 0.12$).

The lack of predictive power of our polygenic score for lottery-based risk tolerance is surprising, as this phenotype is considered the workhorse of risk preference measurement in economic theory¹³⁵. It is important to note, however, that this null result does not necessarily mean that our polygenic score fails to capture variation in economically relevant risk preferences, given its predictive power for risky economic behaviors such as entrepreneurship and financial risk taking. Indeed, the predictive power of lab-based measures for real world behavior has itself come under scrutiny^{172,173}, and lottery-based measures of risk preferences usually have lower predictive power than self-reported measures^{4,6,170,171}. Further, recent research in both economics and psychometrics has found modest levels of correlations between lottery-elicited measures of risk tolerance and the general-risk-tolerance phenotype used in our discovery and replication GWAS^{4,6}. Our results thus add to the small but growing body of evidence that lottery-based and self-reported measures of risk tolerance may indeed capture different aspects of decision-making over uncertainty.

10.3.2 Personality traits

Our results provide interesting evidence regarding the complex and interlacing structure of personality traits and their relationship to general risk tolerance. Genetic predisposition for risk tolerance is predictive of openness to experience, sensation seeking, extraversion, behavioral inhibition, and to a lower extent ADHD. Not surprisingly, these domains are the ones that have been theoretically and phenotypically connected to an individual's behavior in new, unexpected, and uncertain circumstances. Furthermore, these results corroborate our findings of positive

^{ccc} As mentioned above, due to data access limitations, the summary statistics used to construct the score in the STR cohort were based on a meta-analysis that excluded the 23andMe cohort, and therefore had a much smaller GWAS sample size (n_{GWAS} of 466,571).

genetic correlations between general risk tolerance and openness, extraversion, and ADHD (**Supplementary Information section 7.3**).

Looking at the Big Five personality traits, our LDpred-GWAS score is predictive of openness to experience (with incremental R^2 estimates of 1.45% and 0.42% in the HRS and NTR^{fff} cohorts, respectively), extraversion (0.94% in HRS and 0.17% in NTR), and to a lower extent agreeableness (0.09% and 0.11%). The estimated coefficients on the LDpred-GWAS scores for these three phenotypes always have a P value $\leq 0.4\%$. In contrast, conscientiousness was never predicted by our polygenic score, and neuroticism was significantly predicted in the HRS cohort (incremental R^2 of 0.31%), but not in the NTR cohort.

The results described above are broadly consistent with the estimated sign and magnitude of the genetic correlations between general risk tolerance and extraversion, openness, and neuroticism (**Supplementary Information section 7.3**).

Finally, our LDpred-GWAS score was predictive of sensation seeking (with incremental R^2 estimates of 1.13% for the composite sensation seeking phenotype, and ranging from 0.33% to 0.75% for the four sensation seeking subscales), behavioral inhibition (0.50%), and ADHD (0.50%). Furthermore, the estimates of the coefficients on the scores are all highly significant and positive, as expected. No score is predictive of locus of control. Using the summary statistics from the MTAG analysis does not improve our predictive power substantially.

10.3.3 Risky behaviors

(1) Main prediction results

Supplementary Table 10.3A reports the predictive power of the scores for risky behaviors, while **Supplementary Table 10.3B** reports the predictive power of the scores for precautionary behaviors.

Overall across all our validation cohorts, the predictive power of the LDpred-GWAS scores is low but positive across a wide range of risky and precautionary behaviors, with incremental R^2 estimates ranging from 0.02% to 1.36%. Although most incremental R^2 estimates are low, the estimated coefficients on the LDpred-GWAS score have the expected sign for 22 of the 25 regressions we ran ($P = 8 \times 10^{-5}$), and are always in the expected direction whenever the estimated coefficient is significant at the 5% level (16 of the 25 measures, $P = 2 \times 10^{-15}$)^{ggg}.

Our LDpred-GWAS score are predictive of economic behaviors such as being an entrepreneur (1.36%) or owning a business (0.57%), as well as health behaviors such as number of sexual partners (0.65% and 0.79% in Add Health and UKB-siblings), automobile speeding propensity (0.59%), ever using cannabis (0.37%), smoking (incremental R^2 ranging from 0.10% to 0.25% depending on cohort and phenotype), drinking (incremental R^2 ranging from 0.02% to 0.19%, depending on cohort and phenotype), and the first PC of risky behaviors (0.96%). For precautionary behaviors, both life and health insurance coverage are significantly predicted by our LDpred-GWAS scores, with an incremental R^2 of 0.17% and 0.10%, respectively. In contrast, our

^{fff} As mentioned above, the scores in the NTR cohort were also based on a meta-analysis that excluded the 23andMe cohort.

^{ggg} These P values refer to the probability of finding this many concordant signs or significant coefficients; they are derived from one-tailed binomial tests (the first coming from 25 coin flips with probability of 0.5; the second from 25 coin flips with probability of 0.05).

scores are not significantly predictive of financial market participation, equity share or share of financial asset, age of smoking initiation, fried food consumption, and preventative health care utilization.

Several results stand out. The estimated coefficient imply that a one-standard-deviation increase in the polygenic score is associated with a 1.6 percentage points higher probability of being an entrepreneur, 3.4 percentage points higher probability of having owned a business, 2.2 additional lifetime sexual partners, 3.5 percentage points higher probability of ever having smoked marijuana, and a 2 to 3 percentage points higher probability of ever being a smoker.

Using the MTAG summary statistics to construct the LDpred scores in the Add Health, HRS, and UKB-siblings cohorts generally increases our predictive power. For example, the incremental R^2 estimates for the first PC of risky behaviors and for number of sexual partners double, to 1.96% and 1.53% respectively. In the case of ever using cannabis, the incremental R^2 increases by ~0.40% relative to using the LDpred-GWAS and Classical-GWAS scores.

(2) Robustness to inclusion of additional control variables

To further assess the potential for using polygenic scores in empirical research, we investigated whether the predictive power of our polygenic scores is robust to including in the regressions additional control variables for cognitive performance, personality traits, and educational attainment. We obtained very similar point estimates but larger confidence intervals than in the regressions that do not include these additional variables. **Supplementary Table 10.4** displays these results. Note that, due to missing observations in the additional control variables, the sample sizes decrease (in particular, the samples for the STR cohort includes males only, as the cognitive performance measure is only available for males). The pattern of results is nonetheless very similar, though the confidence interval of the estimated coefficients and incremental R^2 values generally increase, reducing our ability to distinguish them from zero. Above and beyond the measures of personal characteristics, our polygenic scores of risk tolerance are still significantly predictive of being an entrepreneur, owning a business, the first PC of risky behaviors, the number of sexual partners, automobile speeding propensity, ever using marijuana, ever being a smoker, being an excessive drinker and drinks per week (in the STR and UKB-siblings cohorts only), and having health or life insurance. Just as for the results displayed in **Supplementary Table 10.3**, LDpred scores constructed using either the GWAS or the MTAG summary statistics are usually more predictive than Classical-GWAS scores.

Furthermore, the predictive power of our polygenic scores for many risky behaviors is comparable in magnitude to that of cognitive performance, personality, and educational attainment. The estimated 95% confidence intervals of the incremental R^2 of our polygenic scores almost always contain the estimated incremental R^2 of cognitive performance, educational attainment, or personality traits.

10.3.4 Height as a negative control

Neither the LDpred-GWAS nor the Classical-GWAS scores significantly predict height in the STR cohort (**Supplementary Table 10.1**). In the UKB-siblings cohort, the incremental R^2 are indistinguishable from zero but, given the large sample size, the coefficients of the LDpred-GWAS and LDpred-MTAG scores are statistically different from zero. These observations, combined with

the above results, suggest that our scores capture some true polygenic signal for general risk tolerance and that our above results are not artifacts of our estimation procedure or of chance.

10.3.5 Concerns about overfitting in the prediction analysis in the UKB-siblings cohort

As indicated above in **Supplementary Note section 10.2**, polygenic scores for the prediction analysis in the UKB-siblings cohort were constructed using summary statistics from a meta-analysis of the 23andMe cohort and a subset of the UKB that excluded all individuals in the UKB-siblings cohort. We will henceforth refer to that subset as the “UKB-nonsibs” cohort.

Although no single individual is present in both the UKB-siblings cohort and in the data used for the meta-analysis, some individuals in the UKB-siblings cohort have relatives in the UKB-nonsibs cohort, and this could lead to overfitting⁹⁵. We determined that the UKB-siblings cohort includes 798 individuals who are ~50% related to an individual in our UKB-nonsibs sample (these likely are parent-child pairs); 1,836 individuals who are ~25% related to an individual in our UKB-nonsibs sample (e.g., half-siblings, or aunt-nephew pairs); and 8,738 individuals who are ~12.5% related to an individual in our UKB-nonsibs sample (e.g., third-degree relatives such as first cousins).

To assess the degree of effective sample overlap across the UKB-siblings and the UKB-nonsibs cohorts, we estimated the intercept in a bivariate LD Score regression²⁴ using the summary statistics of GWAS of general risk tolerance in the UKB-siblings and in the UKB-nonsibs cohorts.

According to theory, the intercept is equal to $\rho N_S / \sqrt{N_1 N_2}$, where ρ is the phenotypic correlation among the N_S individuals included in both samples (so here, $\rho = 1$), and N_1 and N_2 are the sample sizes of the UKB-siblings and the UKB-nonsibs cohorts. From this, we obtained an estimate of N_S : $\hat{N}_S = 837$ ($SE = 654$). This estimate is not significantly different from zero.^{hhh}

Next, to verify that sample overlap does not substantially bias the results of our UKB-siblings prediction analysis, we reran that analysis for general risk tolerance after excluding all individuals in the UKB-siblings cohort with any third-degree or higher relative in the UKB-nonsibs cohort. The results were very similar to those of our baseline analysis: we estimated an incremental R^2 of 1.55% (CI: 1.26% - 1.87%). By comparison, our baseline estimate of the incremental R^2 is 1.62% (CI: 1.37% - 1.90%). We thus conclude that overfitting is unlikely to be a concern for our UKB-siblings prediction analysis.

10.4 Expected predictive power of general-risk-tolerance polygenic score

As mentioned before, the predictive power of our polygenic scores is within the range expected according to theory, when cross-cohort heterogeneity of the GWAS and predicted phenotypes’ SNP heritabilities are taken into account. Daetwyler *et al.*¹³² have developed a theoretical formula for the expected predictive power of a polygenic score, when predicting the phenotype whose

^{hhh} At first glance, this value of N_S seems low relative to the 13,448 UKB-siblings individuals who are related to someone in the UKB-nonsibs cohort. However, we note that the effective sample overlap induced by individuals with relatives depends on both (1) the individuals’ genetic relatedness, weighted by the heritability of risk tolerance, and (2) the correlation between the non-genetic components of their risk tolerance, weighted by one minus the heritability of risk tolerance. The correlation between the non-genetic components is likely much lower than the genetic relatedness, especially for second-degree relatives who are unlikely to cohabitate and to be exposed to highly correlated environmental influences. This depresses the effective sample overlap, relative to what one would expect naively based on a weighted sum of the degrees of genetic relatedness.

GWAS summary statistics were used to construct the score in an independent cohort. The formula assumes independent SNPs, equal heritability of the phenotype in the discovery and validation cohorts, and perfect genetic correlation of the phenotype across cohorts. Daetwyler *et al.*¹³² show that, under those assumptions, the expected predictive power of a polygenic score is equal to:

$$E(R^2) = h^2 \left(\frac{1}{1 + 1/\lambda h^2} \right)$$

where h^2 is the SNP heritability of the phenotype (which is assumed to be the same in the validation cohort and in the GWAS whose summary statistics are used to construct the score), and $\lambda = n_{GWAS}/M$ is the ratio of the sample size in the discovery GWAS (n_{GWAS}) and the effective number of SNPs evaluated in the validation cohort (M). For the discovery meta-analysis we use in our Add Health ($n_{GWAS}=975,353$) and UKB-siblings ($n_{GWAS}=937,353$) analyses, h^2 is 0.041 when estimated by LD Score regression (**Supplementary Table 4.1**) and 0.045 when estimated using HESS (**Table 1**); in the STR cohort ($n_{GWAS} = 458,558$), h^2 is 0.055 when estimated by LD Score regression and 0.063 when estimated by HESS (**Supplementary Table 6.1**)ⁱⁱⁱ; and a reasonable value for M is 60,000, which is the midrange of 50,000 to 70,000 suggested by Wray *et al.* 2013⁹⁵.

Given a range of h^2 values between 0.041 and 0.045 (0.055 and 0.063 for the STR cohort), this formula suggests an expected predictive power between 1.60% to 1.86% in the UKB-siblings cohort, 1.64% to 1.90% in the Add Health cohort, and between 1.63% and 2.05% in the STR cohort. Our estimated incremental R^2 's for general risk tolerance are 1.62% (CI: 1.37% – 1.90%) in the UKB-siblings cohort and ~1.0% in both the Add Health and STR cohorts—slightly less than what we would expect given the Daetwyler formula.

One explanation for the relative underperformance of our polygenic scores is cross-cohort heterogeneity. To account for this, De Vlaming *et al.*¹³³ generalize the Daetwyler formula by allowing for unequal heritability between the prediction (h_p^2) and discovery cohorts (h_D^2) and imperfect genetic correlation ($r_g < 1$), while still assuming independent SNPs. With these generalizations, the expected predictive power of a polygenic score becomes:

$$E(R^2) = r_g^2 h_p^2 \left(\frac{1}{1 + 1/\lambda h_D^2} \right).$$

As before, we let h_p^2 and h_D^2 range between 0.041 and 0.045 (0.055 and 0.063 for the STR cohort). For the UKB-siblings cohort, we estimate a genetic correlation of 0.93 between the discovery and validation cohorts (using the full UKB GWAS instead of a GWAS of the UKB-siblings subset to increase precision of this estimate). For the Add Health and STR cohorts, where sample sizes are too low to permit direct estimation of the genetic correlation, we consider r_g values between 0.7 and 0.9.

The expected incremental R^2 now ranges between 1.51% and 1.61% for the UKB-siblings cohort, 0.80% to 1.54% for the Add Health cohort, and between 0.80% and 1.66% for the STR cohort. The observed predictive power of our score is now consistent with what we would expect in theory.

ⁱⁱⁱ These heritability estimates for the STR cohort are those for the UKB cohort only (and not those for the meta-analysis of the UKB and replication cohorts whose summary statistics were used to construct the scores in the STR cohort). These estimates are higher because they are not attenuated by cross-cohort heterogeneity, unlike the heritability estimates for the GWAS whose summary statistics were used in the Add Health and UKB-siblings analyses.

This demonstrates the importance of considering cross-cohort heterogeneity when forming expectations of the performance of polygenic scores.

De Vlaming *et al.*¹³³ additionally allows for imperfect genetic correlations between individual cohorts within the discovery sample. When we account for this within-discovery sample heterogeneity (using the estimated genetic correlations reported in **Supplementary Information section 7.4.2**, which range from 0.76 to 0.83), our expected R^2 ranges become 1.44%% – 1.67% for the UKB-siblings cohort, 0.84% – 1.62% for the Add Health cohort, and 0.80% – 1.67% for the STR cohort. These ranges remain consistent with our observed R^2 estimates.ⁱⁱⁱ

Lastly, our polygenic score performs consistently better in the UKB-siblings validation cohort than in the Add Health and STR validation cohorts, given our expected R^2 values. This discrepancy could be explained by cross-cohort heterogeneity that we were unable to account for. For example, we have assumed throughout that $h_p^2 = h_D^2$; however, the heritability of risk tolerance may be systematically lower in the Add Health and STR cohorts, where direct heritability estimation was not possible.

Thus, while polygenic scores constructed from our GWAS results can already aid the study of some related phenotypes, future research would benefit from the collection of measures of risk tolerance that are more similar across cohorts.

ⁱⁱⁱ We do not account for imperfect genetic correlations between cohorts within our replication meta-analysis; these cohorts only account for 3.5% of the individuals in our discovery meta-analysis and are unlikely to affect our results.

10.5 Appendix: definition of the income risk gamble variable in the HRS cohort and merger of the STR1 and STR2 cohorts

10.5.1 Income gamble risk tolerance in the HRS cohort

In the HRS cohort the income gamble risk question was slightly modified after the first wave¹³⁴. In the first wave, respondents were first asked the following question:

“Suppose that you are the only income earner in the family, and you have a good job guaranteed to give you your current (family) income every year for life. You are given the opportunity to take a new and equally good job, with a 50-50 chance it will double your (family) income and a 50-50 chance that it will cut your (family) income by a third. Would you take the new job?”

Respondents who answered “yes” to the first question were asked this follow-up question:

“Suppose the chances were 50-50 that it would double your (family) income, and 50-50 that it would cut it in half. Would you still take the new job?”

Respondents who answered “no” to the first question were asked this follow-up question:

“Suppose the chances were 50-50 that it would double your (family) income and 50-50 that it would cut it by 20 percent. Would you then take the new job?”

In the original variable coding, the variable was coded as 1 if the respondent accepted the new job with the risk of cutting income by half, and a value of 4 if the respondent always chose to stay with the job with a guaranteed income. We reverse coded the original phenotype so that a higher value implies higher risk tolerance.

The income gamble risk tolerance question was not asked in waves 2 and 3. In wave 4 and all subsequent waves, the initial question was:

“Suppose that you are the only income earner in the family. Your doctor recommends that you move because of allergies, and you have to choose between two possible jobs. The first would guarantee your current total family income for life. The second is possibly better paying, but the income is also less certain. There is a 50-50 chance the second job would double your total lifetime income and a 50-50 chance that it would cut it by a third. Which job would you take -- the first job or the second job?”

If the first job was chosen as the answer to the initial question, then this follow-up question was asked:

“Suppose the chances were 50-50 that the second job would double your lifetime income and 50-50 that it would cut it by twenty percent. Would you take the first job or the second job?”

If the first job was chosen as the answer to the follow-up question, then a second follow-up question was asked:

“Suppose the chances were 50-50 that the second job would double your lifetime income and 50-50 that it would cut it by 10 percent. Would you take the first job or the second job?”

If instead the second job was chosen as the answer to the initial question, then a different follow-up question was asked:

“Suppose the chances were 50-50 that the second job would double your lifetime income, and 50-50 that it would cut it in half. Would you take the first job or the second job?”

If the first job was chosen as the answer to the follow-up question, then a second follow-up question was asked:

“Suppose the chances were 50-50 that the second job would double your lifetime income and 50-50 that it would cut it by seventy-five percent. Would you take the first job or the second job?”

In the original variable coding, the respondent is given a value of 1 if the respondent accepts the second job with the risk of cutting the income by seventy-five percent, and a value of 6 if the respondent always chooses to stay with the job with a guaranteed income. Again, we reverse coded the responses so that a higher value corresponds to a higher risk tolerance. Further, since this survey measure has two additional response categories relative to the measure used in the first wave, we converted this survey measure’s responses to the 4-point scale that disregards the second follow-up questions; a value of 5 or 6 corresponds to 4, a value of 4 corresponds to 3, a value of 3 corresponds to 2, and value 1 or 2 corresponds to 1.

We constructed the income gamble risk tolerance phenotype as the average response across waves, as follows. We first computed the residuals for wave w as

$$\begin{aligned} y_{w,i} &= \mathbf{X}_i \boldsymbol{\beta}_w + \epsilon_{w,i}, \\ \hat{\epsilon}_{w,i} &= y_{w,i} - \mathbf{X}_i \hat{\boldsymbol{\beta}}_w \end{aligned} \quad (10)$$

where $y_{w,i}$ is income gamble risk tolerance in wave w for individual i and \mathbf{X}_i contains the intercept and the control variables (birth year, birth year squared, birth year cubed, sex, as well as three interaction terms between sex and the three birth year variables). We then calculated the average residual across waves as $\hat{\epsilon}_i = \overline{\hat{\epsilon}_{w,i}}$. To ensure this averaged residual is not associated with any of the control variables in \mathbf{X}_i either, we computed residual p_i as

$$\begin{aligned} \hat{\epsilon}_i &= \mathbf{X}_i \boldsymbol{\beta} + \eta_i, \\ p_i &= \hat{\epsilon}_i - \mathbf{X}_i \hat{\boldsymbol{\beta}}, \end{aligned} \quad (11)$$

and used this residual p_i is used as the phenotype for the prediction in the HRS cohort. The sample size for income gamble risk tolerance is 7,302 in the HRS cohort (**Supplementary Table 10.1**).

10.5.2 Merging the STR1 and STR2 cohorts

Since the STR1 and STR2 cohorts have been genotyped with different genotyping arrays (for details, see **Supplementary Table 2.1**), we kept them separate for the GWAS of general risk tolerance (**Supplementary Information section 2**). However, we merged the cohorts for the prediction analyses. First, we constructed and standardized the polygenic scores and calculated the 10 genetic principal components (PCs) separately in the STR1 and STR2 cohorts. We then merged the individual-level data by pooling the polygenic scores into a single variable. Two separate sets of PCs were used, imputing the value 0 for the 10 PCs of STR2 for the individuals in the STR1 cohort, and vice versa. In all the prediction analyses in the merged STR cohort, we included a dummy variable indicating whether the individual belonged to the STR1 or STR2 cohort.

11 Biological annotation: testing hypotheses about specific genes and gene sets

Earlier studies have shown that risk tolerance and related individual characteristics are moderately heritable^{7,174}, thus providing a motivation to investigate the biological and molecular bases of this heritability. Indeed, there is a voluminous literature that links specific biological pathways (i.e. brain regions, neuronal populations, hormones, receptors, and neurotransmitters) to decision making and behavior. Different research designs and proxies for biological pathways have been used, all with specific advantages and disadvantages (see, for example, Nave *et al.* (2015)¹⁷⁵ for a review of studies on the role of oxytocin on trust).

One part of this literature is candidate gene studies that have attempted to leverage insights from psychology and biology to derive hypotheses that could be tested using specific genes as proxies for biological pathways. Although the hypothesis-based approach seems intuitively reasonable, it is now widely accepted that findings from candidate gene studies on human behavior often fail to replicate. The reasons for this include very small effects of common genetic variants on human behavior, small sample sizes in candidate gene studies that led to underpowered statistical tests, publication bias, and insufficient controls for correlations between genotypes and relevant environmental conditions^{15,16,18,20,176–179}.

We systematically reviewed the prior literature on biological pathways that have been hypothesized to be linked to risk tolerance. Then, we used MAGMA¹⁸⁰ on the summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance ($n = 975,353$; as usual, we applied genomic control using the intercept of the LD Score regression to these summary statistics prior to the analyses) to re-evaluate the findings of this literature.

In addition, since risk tolerance has been hypothesized to be under recent evolutionary pressure^{181–183}, we tested whether a set of genes previously identified as having been under evolutionary pressure in Europeans is associated with general risk tolerance.

11.1 Literature review

11.1.1 Search algorithm

We conducted a comprehensive literature review on biological pathways that may influence risk tolerance. Consistent with our GWAS, we restricted our review to research involving healthy individuals. Thus, we excluded studies that focused on neuropathologies, addictions, or other disorders. However, we included articles using animal models to study risk tolerance because the investigated biological mechanisms may have similar effects on human behavior. Our search algorithm employed a relatively broad definition of risk tolerance and included psychometric as well as behavioral measures (e.g. self-reported risk tolerance, choices between monetary lotteries, or gambling tasks). Risk tolerance among animals was typically measured by observing choices between probabilistic food options or by quantifying risk-assessment behavior during exploration. Following the analysis plan for our GWAS of general risk tolerance, we considered impulsivity and novelty-seeking as conceptually different traits and excluded them from this literature review. We included all proxies of biological pathways that are tested in the literature, including candidate genes, molecules represented by specific receptors, pharmacological interventions that are used to

manipulate specific pathways (e.g., nasal oxytocin spray, neurotransmitter receptor blockers, agonists, antagonists), as well as very distal proxies like the 2D:4D digit ratio.

We employed two parallel approaches to translate these criteria into search algorithms: first, we used a “bottom-up” approach and searched for studies investigating the relationship between risk taking and any biological pathway. The “bottom-up” search criterion we used was:

(“risky behavior” OR “risk-aversion” OR “risk preference” OR “risk-taking” OR “risk-seeking”) AND (“GWAS” OR “gene” OR “SNP” OR “allele” OR “genetic” OR “candidate” OR “hormones” OR “neurotransmitter” OR “neuropeptide” OR “biology” OR “biological pathway”).

We also conducted a “top-down” search which specifically cued our search criterion with biological pathways that were mentioned in the prior literature on risk tolerance. This “top-down” search criterion was:

(“risky behavior” OR “risk-aversion” OR “risk preference” OR “risk-taking” OR “risk-seeking”) AND (“catecholamines” OR “monoamines” OR “dopamine” OR “adrenaline” OR “noradrenaline” OR “norepinephrine” OR “glutamate” OR “serotonin” OR “GABA” OR “BDNF” OR “testosterone” OR “acetylcholine” OR “NMDA” OR “AMPA” OR “testosterone” OR “estradiol” OR “progesterone” OR “cortisol” OR “glucocorticoid” OR “vasopressin” OR “oxytocin” OR “ghrelin” OR “leptin”).

We used both of these algorithms in two different search engines; the ISI Web of Knowledge and Google Scholar. Both search engines yielded highly overlapping results.

We screened approximately 1,000 articles and found 132 which matched our criteria, spanning different scientific fields (from management to neuroscience), different models (e.g. human or rodent), different risk measures (from psychometric to behavioral measures), different biological pathways (e.g. monoamines, sex hormones), and different approaches (from genotyping to pharmacological manipulations). We included articles regardless of whether the reported results were statistically significant or not.

11.1.2 Results of the literature review

The results of our literature review are compiled in **Supplementary Table 11.1**. Overall, the literature on biological mechanisms for risk tolerance is relatively recent: most studies were published after 2003. However, earlier work on related concepts that were excluded by our selection criteria (such as novelty seeking) exists.

Three main biological mechanisms comprising five biological pathways have been tested by this literature. The first of these is the relationship between risk tolerance and sex hormones, especially testosterone and estrogens. Testosterone and estrogens are the two primary steroid sex hormones in humans. They are principally (but not exclusively) synthesized in the ovaries (estrogen) and testicles (testosterone), and are responsible for the development and regulation of the male (testosterone) and female (estrogen) reproductive system and secondary sex characteristics. The link between those hormones and risk taking is generally motivated by (and discussed as

corroborating of) the observed behavioral and attitudinal differences between males and females. A large part of the literature testing the relationship between risk tolerance and sex hormones uses the distal 2D:4D digit ratio to approximate prenatal testosterone levels. Some studies use measures of sex hormones that are derived from saliva or blood samples.

The second frequently examined biological mechanism is the relationship with the neurotransmitters dopamine and serotonin. Those two monoamines have been repeatedly implicated in behavior and decision-making: dopamine plays a major role in reward-motivated behavior and in addiction, whereas serotonin is implicated in mood. Studies testing the associations between these monoamine pathways and risk tolerance typically test genetic variants implicated in the synthesis or the signaling pathways of these neurotransmitters (e.g., *DRD4*, *SERT* or *5-HTTLPR*). Alternatively, pharmacological manipulations are used including blockers, agonists, or antagonists of specific dopamine or serotonin receptors.

The third main biological pathway tested by the literature involves cortisol. Cortisol is another steroid hormone belonging to the glucocorticoid class, and is notably produced in response to stress. Cortisol triggers gluconeogenesis (the formation of glucose), and activates anti-stress and anti-inflammatory pathways. Studies testing the associations between the cortisol pathways and risk tolerance typically use a biochemical quantification of cortisol level in salivary or blood samples as an independent measure.

In sum, our literature review highlighted five main biological mechanisms potentially underpinning the risk-tolerance phenotype: the testosterone, estrogen, dopamine, serotonin and cortisol pathways.

11.2 Gene analysis and competitive gene-set analyses with MAGMA

We used MAGMA¹⁸⁰ together with the summary statistics from the meta-analysis of the discovery and replication GWAS of self-reported general risk tolerance to re-evaluate the findings of this literature.

We conducted several types of analyses based on these results. First, for all five mechanisms we constructed gene sets, based on external databases of biological function, and tested these gene sets for association with risk tolerance. Second, for two of those pathways (dopamine and serotonin), there are a significant number of candidate gene studies in humans. These candidate gene studies most commonly test a specific subset of genes and genetic variants (SNPs or structural variants) within the dopamine and serotonin pathways. We therefore built a gene set from 15 of the 17 most commonly tested genes of those two pathways (**Supplementary Table 11.4**) in order to re-evaluate the associations of those candidate genes with risk tolerance. Third, we tested specific SNPs in these 15 genes that have been tested in the prior candidate gene literature. Lastly, we tested whether a set of genes that has been identified as having been under selection among Europeans was enriched for association with general risk tolerance.

MAGMA can perform two types of analyses: a gene analysis and a competitive gene-set analysis. For a given gene, a MAGMA gene analysis tests the joint association of all the SNPs in the gene with the phenotype. This aggregation of the SNPs reduces the number of tests compared to a SNP-based analysis, which results in an increase in statistical power and may lead to the detection of joint effects of multiple weaker SNP associations that would otherwise be missed. A disadvantage

of gene analysis is that it is uninformative about the direction of the effect, as it simultaneously tests multiple SNPs that can have opposite effects within a gene.

For a given gene set, a MAGMA competitive gene-set analysis tests whether the genes in the gene set are more strongly associated with general risk tolerance than the other genes in the genome. Gene sets are groups of genes that share certain characteristics (e.g., biological or functional characteristics). Like a gene analysis, a competitive gene-set analysis has increased statistical power relative to a SNP-based analysis, as it involves fewer tests. Moreover, a gene-set analysis allows tests of specific biological hypotheses by defining appropriate gene-sets, and may provide direct insights into the underlying biological pathways or cellular functions of the phenotype. MAGMA uses a regression framework that compares the mean association of the genes in the gene set with the mean association of all other genes in the genome (equivalent to a one-sided two-sample *t*-test). To account for possible LD between genes, MAGMA uses a gene correlation matrix, which was based on the European ancestry samples from the 1000 Genomes project phase 1⁹¹. Additional covariates are included in the regression to correct for any confounding effects of gene size and gene density (gene size, log gene size, gene density, log gene density).

11.2.1 Competitive gene-set analysis with MAGMA: Testing gene sets related to dopamine, serotonin, testosterone, estrogen, and glucocorticoids

As mentioned above, not all of the biological pathways that were studied in the previous literature on risk tolerance were studied using candidate genes as proxies for these pathways. Furthermore, the candidate genes (e.g. *DRD4*) that were studied are not the only genes that are involved in the respective biological pathways (e.g. dopamine). Thus, to begin with, we constructed and tested gene sets that represent the five major biological pathways that were previously studied in the context of risk tolerance. (We note that we also performed *ex post* MAGMA gene set analyses for GABA and glutamate neurotransmitters. We describe these analyses in **Supplementary Information section 12.5**.)

We selected all gene sets related to the five major biological pathways (dopamine, serotonin, testosterone, estrogen and cortisol) from the Molecular Signature Database (MSigDB, v5.1)¹⁸⁴, and from the gene sets compiled by Hawrylycz *et al.* (*Nature Neuroscience* 2015)¹⁸⁵. There were 38 gene sets related to the five pathways, from which we removed four duplicate gene sets. The remaining 34 gene sets had much overlap within each biological pathway, and we therefore merged all the gene sets belonging to a given pathway, resulting in five gene sets corresponding to each of the five pathways (dopamine, serotonin, testosterone, estrogen and glucocorticoids (cortisol)). We report both the initial and the merged gene sets in **Supplementary Table 11.2**.

We conducted a competitive gene-set analysis with MAGMA to test each of these five gene sets for association with risk tolerance. We applied a resampling-based *P* value adjustment¹⁸⁶ using 10,000 permutations to correct for multiple testing (as we conducted a test for each of the five gene sets).

None of these five gene sets showed a significant association with general risk tolerance after correction for multiple testing (the lowest corrected *P* value was 0.27, **Supplementary Table 11.3**).

11.2.2 Gene analysis and competitive gene-set analysis with MAGMA: Testing candidate genes from the prior literature

We used MAGMA to conduct a gene analysis to test the candidate genes identified in our literature review for association with general risk tolerance, employing all of the SNPs in our GWAS summary statistics within the physical location of the genes being tested. 15 of the 17 candidate genes that we identified in the literature review were autosomal and were thus eligible for testing (since our GWAS of general risk tolerance was restricted to autosomal SNPs). In addition to the gene analysis, we conducted a competitive gene-set analysis with MAGMA to test whether the genes in the gene set comprising these 15 genes are more strongly associated with general risk tolerance than the other genes in the genome.

From the gene analysis, none of the genes showed a significant association with general risk tolerance after Bonferroni correction to correct for multiple testing (the 15 genes are not in LD; **Supplementary Table 11.4**). From the competitive gene-set analysis, the set of 15 candidate genes was also not significantly associated with risk tolerance ($P = 0.55$, **Supplementary Table 11.4**).

Our MAGMA competitive gene-set analysis of the cortisol, dopamine, serotonin, estrogen, and testosterone gene sets has some likely limitations. First, the statistical power of MAGMA competitive gene-set analyses barely increases with increasing sample sizes¹⁸⁷. Second, important effects of numerous single SNPs in a gene or gene set may be overshadowed by the high average P values of other SNPs in the gene or gene set. Third, only specific pathways within the tested gene sets might be relevant, and merging many pathways into major cortisol, dopamine, serotonin, estrogen, and testosterone gene sets decreased our statistical power to discover specific pathways. In **Supplementary Information section 12.7.4**, we further discuss these limitations in the context of our *ex post* MAGMA competitive gene-set analysis of four glutamate and GABA gene sets; that analysis also returned null results, despite the fact that our Gene Network and DEPICT analyses both point to a role for glutamate and GABA neurotransmitters.

Importantly, however—and unlike for glutamate and GABA—none of the bioinformatics analyses we report in **Supplementary Information section 12** point to the cortisol, dopamine, serotonin, estrogen, and testosterone pathways or related genes either. Further, as reported above none of the 15 commonly-tested candidate genes were significant in our MAGMA gene analysis after Bonferroni correction for 15 tests; by contrast, our MAGMA gene analysis of ~18,000 genes identified several glutamate and GABA genes that were significant after Bonferroni correction for ~18,000 tests (**Supplementary Information section 12.2**). Moreover, as we discuss in **Supplementary Information section 11.2.3** just below, none of the SNPs tagging the 15 most commonly-tested autosomal candidate genes within the dopamine and serotonin pathways have non-negligible effects on general risk tolerance. Therefore, while our MAGMA competitive gene-set analysis may suffer from some limitations, other analyses we conducted also fail to point to a role for the cortisol, dopamine, serotonin, estrogen, and testosterone pathways, and instead point to a role for glutamate and GABA neurotransmitters.

11.2.3 Replication of specific SNPs tested in the prior candidate gene literature

As a complement to the MAGMA gene-based tests, we also tested the individual SNPs that have been tested for association with risk tolerance in the previous literature and that are located within the 15 autosomal genes tested with MAGMA. We used the summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance for these tests. Out of

the 15 autosomal genes, three have been tested in the previous literature for association with alleles of an underlying structural variant, and we treat these separately below, where we looked up a total of 18 SNPs available in our GWAS summary statistics for these three genes. The remaining 12 autosomal genes contained 75 SNPs that were individually tested in the previous literature, of which 74 were available in our meta-analysis of the discovery and replication GWAS of general risk tolerance. Thus, our GWAS summary statistics contain the majority of candidate SNPs within these genes that were tested in the previous literature, and we performed Bonferroni-correction for testing 92 candidate SNPs.

Out of the 74 previously tested SNPs within the 12 autosomal genes, two replicated with Bonferroni-corrected P values less than 0.05. The SNPs were rs36339 (in the gene *SLC1802*; Bonferroni-corrected $P = 0.004$), and rs174699 (in the gene *COMT*; Bonferroni-corrected $P = 0.034$). The R^2 's of rs36339 and rs174699 in the summary statistics of our discovery GWAS of general risk tolerance are 0.00185% and 0.00141%, respectively. By comparison, the smallest R^2 among the lead SNPs from our discovery GWAS of general risk tolerance is almost twice as large, at 0.0032% (rs1935571). The effects of rs36339 and rs174699 are considerably smaller than what could be found at the genome-wide significance level with our current discovery sample of 939,908 individuals, and they are several orders of magnitude smaller than what candidate gene studies in the existing literature—with typical sample sizes rarely exceeding a few hundred to a few thousand individuals—could have found. The sample sizes required to have 80% statistical power to detect the effects of rs36339 and rs174699 at the 5% level of significance (without Bonferroni correction) would be 424,260 and 556,654, respectively; the corresponding sample sizes at the genome-wide significance level (5×10^{-8}) would be 2,140,573 and 2,808,560, respectively. We therefore found that 10 of the 12 hypothesized autosomal candidate genes harbor no candidate SNPs that replicate, while two (*SLC1802* and *COMT*) each contain a SNP that is suggestively associated with general risk tolerance (with a P value less than 0.05 after Bonferroni correction for 92 tests) but whose effect is very small.

In the previous literature, three of the 15 autosomal candidate genes (*DRD4*, *DRD5* and *SLC6A4* (the latter is also commonly referred to as *SERT* or *5-HTT*)) have mainly been tested for association with behavioral traits using alleles of structural variants located within these genes. For example, risk tolerance has previously been tested for association with a variable number tandem repeat in the *DRD4* gene referred to as the 7R polymorphism¹¹. Our GWAS summary statistics include nine SNPs within the *DRD4* gene. Although we do not know the exact LD between these nine SNPs and the 7R polymorphism, it is very likely that at least some of them tag the 7R polymorphism very well because the *DRD4* gene spans only 3,401 base pairs on chromosome 11, and variants in such close physical proximity tend to be in high LD. The lowest Bonferroni-corrected P value of a SNP in the *DRD4* gene is 0.47 (rs201554946). The gene *DRD5* has mainly been tested for association with a microsatellite that can take many different alleles¹⁸⁸, and no SNPs within *DRD5* were found in our literature review to have been tested directly for association with risk tolerance. *DRD5* spans 2,375 base pairs on chromosome 4, in which we have seven SNPs in our GWAS summary statistics, and none of the SNPs have a Bonferroni-corrected P value less than 1. The gene *SLC6A4* has mainly been tested for association with risk tolerance via a degenerate repeat polymorphism in a region of the gene called *5-HTTLPR*¹⁸⁹, and alleles of this repeat polymorphism are tagged by the SNPs rs2129785, rs11867581, and rs25531¹⁹⁰. The SNPs rs2129785 and rs11867581 are available in our GWAS results and their Bonferroni-corrected P values are both 1. Based on these results we cannot reject the null hypothesis that none of these three genes is associated with risk tolerance.

Extended Data Fig. 3.4b and **Fig. 1c** display local Manhattan plots of the areas around the 15 most commonly tested candidate genes in the prior literature on the genetics of risk tolerance. Each local plot shows all SNPs within 500 kb of the gene's borders that are in LD ($r^2 > 0.1$) with a SNP in the gene. As can be seen in **Extended Data Fig. 3.4b**, only one of the 15 areas (around the gene *DRD2*) contains lead SNPs for any of our seven GWAS, and these SNPs are only genome-wide significant in our GWAS of drinks per week. One SNP in the *SLC6A4* (*SERT*) locus is genome-wide significant in our GWAS of adventurousness. However, this SNP (rs112739039) is not a lead SNP in our GWAS of adventurousness because it is located in the locus of another SNP (rs6505239; see **Supplementary Table 3.2**). The nearest gene to rs112739039, *RAB11FIP4*, is located ~1Mb upstream of *SLC6A4*. In our meta-analysis of general risk tolerance, this *SLC6A4* SNP is not genome-wide significant and is included in the clump around rs11080149 with nearest gene *OMG*, which is also located ~1Mb upstream of *SLC6A4* (see **Supplementary Table 3.1**).

The local Manhattan plots of the areas around the 15 candidate genes (**Extended Data Fig. 3.4b** and **Fig. 1c**) can be compared to those of the five main long-range LD regions and candidate inversions described in **Supplementary Information section 3.2** (**Extended Data Fig. 3.4, a and c**, and **Fig. 1, a and b**). These five regions contain numerous lead SNPs for most or all of our seven main GWAS.

In summary, no candidate genes that have been tested with SNPs in the previous literature, nor the candidate genes tested with alleles of structural variation, contain genome-wide significant SNPs in our discovery GWAS, while two candidate genes each contain a SNP with a suggestive association (i.e., with a Bonferroni-corrected P value smaller than 0.05, after Bonferroni correction for 92 tests). However, a sample size of more than 2 million individuals would be required to have 80% power to replicate these two SNPs at the genome-wide significance level, and a sample size of more than 400,000 would be required to replicate these SNPs at the 5% level of significance. We therefore conclude that these two SNPs are relatively unimportant in terms of their effects on general risk tolerance, and that previous candidate gene studies testing these SNPs for association with behavioral phenotypes were severely underpowered to detect their very small effects.

11.2.4 Competitive gene-set analysis with MAGMA: Testing genes under selection among Europeans

A number of theorists have posited that attitudes toward risk have been under evolutionary pressure^{181–183}. In particular, if decisions involving risk have to be made frequently and if decision mistakes have a severe impact on fitness, theoretical models suggest that selective pressure will favor risk attitudes that maximize fitness in a given environment¹⁸². Consequently, changes in the environment can induce selective pressure for different levels of risk tolerance. It is obvious that the environment in which people live and make choices has changed dramatically over the past few millennia, with many fitness-related risk factors declining in importance, and new ones arising (e.g. due to urbanization, technological progress, and increasing populations). Hence, it is conceivable that general risk tolerance has been under selective pressure in our recent evolutionary past.

To test if risk tolerance was subject to recent selective pressure among people of European descent, we constructed a gene-set based on 17 genes that were highlighted as having been under recent selective pressure among Europeans by the 1000 Genomes phase 3 analyses¹⁹¹ (see **Supplementary Table 11.5**). We performed a competitive gene-set analysis with MAGMA and

with the summary statistics from the meta-analysis of our discovery and replication GWAS of general risk tolerance, to test whether the genes in this gene-set are more strongly associated with general risk tolerance than other genes.

The gene-set based on the genes that have been highlighted as having been under recent selective pressure among Europeans showed no association with general risk tolerance (P value = 0.90, **Supplementary Table 11.5**). Thus, this test provides no evidence suggesting that risk tolerance has been subject to recent selective pressure. However, this absence of evidence is not evidence of absence: our test may have failed to detect evolutionary pressure for several reasons, including limited statistical power. Further, we only tested if a set of genes that have been highlighted as having been under evolutionary pressure is associated with general risk tolerance; future research could also test whether genes that are associated with risk tolerance have been under evolutionary pressure.

11.3 Discussion

In summary, we find no evidence of enrichment for the main pathways and genes that had previously been hypothesized to relate to risk tolerance. We also note that none of the bioinformatics analyses we report in **Supplementary Information section 12** point to these pathways or genes either (we note, however, that some brain regions identified in analyses we report below are areas where dopamine and serotonin play important roles.) By contrast, some those bioinformatics analysis point to a role for the glutamate and GABA neurotransmitters.

Our null replication results are consistent with the poor replication record that candidate gene studies in social-science genomics have typically had^{15,16,18,20,176–179}. Given that the sample size of our GWAS is several orders of magnitudes larger than any previous candidate gene study on risk tolerance, our results put a credible, tight upper bound on the effect sizes of the genes that were tested and allow us to rule out the possibility that these genes have particularly large effect sizes compared to typical genetic variants.

However, our results do not imply that the main biological pathways we identified in our literature review are irrelevant for risk tolerance, as variations further downstream in these pathways (e.g. damaged glands, pharmacological interventions) may have much stronger effects on risk tolerance than the genes that were tested.

12 Biological annotation: other bioinformatics analyses

This section reports biological annotation of our GWAS results. Throughout, we focus on our primary phenotype, self-reported general risk tolerance, because it has by far the largest GWAS sample size and because the other main phenotypes analyzed in this paper are genetically correlated with it. The goal of these analyses is to gain biological insight (1) into the genome-wide biological correlates of the general-risk-tolerance phenotype (with a suite of bioinformatic analyses that use GWAS summary statistics for large sets of SNPs across the genome), and (2) about the lead SNPs from our discovery GWAS of general risk tolerance (by looking up the functional status of the lead SNPs and SNPs in LD with them). For all analyses in this section, we use the summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance.

In **Supplementary Information section 12.1**, we use partitioned LD Score regression⁹⁴ to test for polygenic enrichment in specific genomic regions, such as gene transcription start sites, evolutionarily conserved regions, and sections of the genome epigenetically modified in certain tissues.

In **Supplementary Information section 12.2**, we use MAGMA¹⁸⁰ to conduct hypothesis-free analyses to identify specific genes that are associated with general risk tolerance. We also use a co-expression database to gain insight into the functions of the significant MAGMA genes.

In **Supplementary Information section 12.3**, we perform transcriptome-wide analysis with Summary-based Mendelian Randomization (SMR), which leverages genome-wide data to discover genes whose expression significantly associates with general risk tolerance. In addition, the “HEIDI” test feature of SMR is able to discard expression quantitative trait loci (eQTL) associations that are caused merely by linkage, thereby discarding genes that are spuriously associated with general risk tolerance.

In **Supplementary Information section 12.4**, we use DEPICT¹⁹² to prioritize tissues, gene sets, and genes that are implicated by our GWAS results. Like MAGMA, DEPICT is a gene-based tool. However, in contrast to MAGMA, DEPICT uses reconstituted gene sets that are based on co-expression patterns to prioritize genes, gene sets, and pathways.

In **Supplementary Information section 12.5**, to follow up on the results from other biological annotation analyses, we conduct an *ex post* MAGMA gene set analysis¹⁸⁰ to test for enrichment of GABA and glutamate pathways in general risk tolerance.

In **Supplementary Information section 12.6**, we report a number of analyses that annotate our general-risk-tolerance lead SNPs, to gain insights into the biological systems they may affect. First, we check whether our lead SNPs (and SNPs in LD with them) contain protein-altering variants (i.e., genetic variants which result in a structural difference in the protein structure encoded by a gene, collectively known as ‘nonsynonymous’ variants). Second, we look up the lead SNPs (and SNPs in LD with them) in an expression quantitative trait loci (eQTL) database that contains associations between genetic variants and gene expression levels.

Finally, in **Supplementary Information section 12.7**, we highlight the most important results of these analyses and summarize the conclusions we derive from them.

12.1 Functional partitioning of heritability with stratified LD Score regression

12.1.1 Background and methods

In this section, we discuss the results of our stratified LD Score regression analyses⁹⁴. (In addition to ref.⁹⁴, the Supplementary Information of Okbay *et al.*¹⁸ also contains a detailed description of the methodology.) With these analyses, we break down (“partition”) the SNP-based heritability of general risk tolerance across SNPs with various functional genomic annotations. We used the GWAS summary statistics from the meta-analysis of the discovery and replication GWAS of general risk tolerance (after applying genomic control using the intercept of the LD Score regression, as usual) for these analyses.

Stratified LD Score regression is based on the relationship

$$(12) \quad E[\chi_j^2] = N \sum_{c=1}^c \tau_c \ell(j, c) + Na + 1,$$

where $\chi_j^2 = N\hat{\beta}_j^2$ is the GWAS chi-square statistic for SNP j , N is the GWAS sample size, c indexes the functional categories (which do not have to be disjoint), $\ell(j, c)$ is the stratified LD Score of SNP j with respect to functional category c , τ_c is the average contribution to heritability of a SNP due to its membership in category c , and a is a term that measures the contribution of confounding biases such as cryptic relatedness and population stratification.

Finucane *et al.*⁹⁴ present derivations of this equation and show how estimates of τ_c that result from estimating the implied regression can be used to obtain estimates of the heritability ascribable to the various functional categories. Enrichment is then defined as the fraction of the total heritability captured by the functional annotation category divided by the fraction of SNPs in that category.

To partition the SNP-based heritability of risk tolerance using the results of our GWAS meta-analysis, we followed the procedure described by Finucane *et al.*⁹⁴ and applied in two recent papers by Okbay *et al.*^{16,18}. That is, we used the stratified LD scores calculated from the European-ancestry samples in the 1000 Genomes Project (1000G) (accessed on March 14, 2016), but in the regressions themselves included only the chi-square statistics of the ~ 1 million HapMap3 SNPs with minor allele frequency (MAF) > 0.05 . This decision was motivated by the facts that HapMap3 SNPs are a comprehensive set of common genetic markers that can be imputed with high accuracy across different groups and genotyping platforms^{90,167}; that the LD scores of SNPs with low MAFs can introduce too much statistical noise when used in the analysis; and that the per-SNP heritability may substantially differ for common and rare SNPs.

We performed two distinct analyses. We first estimated the stratified LD Score regression for the functional genomic regions of the “baseline” model. The functional categories in the baseline model consist of one category containing all SNPs, 24 categories corresponding to 24 main functional annotations of interest (which include, for example, evolutionarily conserved regions in mammals and regions epigenetically regulated to be accessible to gene transcription), categories corresponding to 500 bp windows around regions belonging to each of these 24 annotations (to correct for spurious associations driven by variants located near the physical borders of the functional genomic regions), and categories corresponding to 100-bp windows around ChIP-seq peaks (regions that are DNase hypersensitive, also known as “open chromatin,” or associated with

the histone marks H3K4me1, H3K4me3, H3K9ac, or H3K27ac). The full baseline model contains 53 predictors (including the predictor for the category containing all SNPs).

Second, to gain tissue-level resolution, we used the tissue-level annotations provided by Finucane *et al.* These consist of 220 cell-type-specific annotations for four histone marks that are subsequently grouped into 10 broad tissue type annotations (Adrenal/Pancreas, Central Nervous System, Cardiovascular, Connective/Bone, Gastrointestinal, Immune/Hematopoietic, Kidney, Liver, Skeletal Muscle, and Other). Histone marks are posttranslational modifications of histones that alter their interaction with the DNA wound around them. For example, the acetylation of lysine (an amino acid present in the histone protein) may facilitate gene expression by reducing the electrical attraction between DNA and the histone, making the DNA strand more receptive for transcription (and thus, gene expression). While certain histone marks are associated with increased DNA transcription, other histone marks' effects on DNA transcription depend on the combination of the chemical form and the location of the modification. For instance, mono-methylation of histone H3K9 (denoted H3K9me1) is associated with activation of gene expression, while tri-methylation of H3K9 (denoted H3K9me3) is associated with repression of gene expression¹⁹³. The association between SNPs and histone modifications differs per tissue and developmental stage, and has been mapped extensively by the RoadMap Epigenomics project¹⁹⁴. For instance, SNPs important for eye function are likely to be “tightly bound” to histone proteins in the liver, where they do not need to be expressed, but “loosely” bound to histones in the eye, where their expression is crucial. Here, we used LDSC partitioning of heritability to test if risk tolerance SNPs are enriched in (or near) tissue-specific histone marks, which would suggest that those tissues might be of biological importance for general risk tolerance.

The functional annotation categories we used here are associated with the histone marks H3K4me1, H3K4me3, H3K9ac and H3K27ac, which are all known to increase transcription rate¹⁹⁵. However, we note that the probability of any gene being transcribed and thus expressed further depends on the presence of other epigenetic modifications, such as epigenetic modifications of the DNA sequence itself (for example, methylation of CpG-islands). This means that the four histone modifications we study here do not deterministically increase gene expression rates, but rather do so in a probabilistic manner.

We added each of the 10 tissue annotations to the baseline model (resulting in 10 separate regressions, each with 54 predictors), and assessed the magnitude and statistical significance of the observed enrichment. To benchmark these tissue-level results, we compared them to the corresponding estimates we obtained using the summary statistics of the most recent GWAS of height¹⁰⁵ (<http://www.broadinstitute.org/collaboration/giant/index.php>).

To correct for multiple hypothesis testing, we applied a Bonferroni correction for 52 two-sided tests in the baseline model (i.e., for 52 annotations^{kkk}), and for 10 two-sided tests in the tissue type models (i.e., for 10 tissue types).

12.1.2 Results: The “baseline” model

The results for the baseline model are shown in **Supplementary Table 12.1**. The baseline annotations “Conserved” and “H3K9ac peaks” are the most enriched, with enrichment estimates that remain significant after Bonferroni correction for 52 tests. The “Conserved” category shows

^{kkk} The baseline model has 53 predictors, but we do not adjust for the predictor for the category containing all SNPs.

the strongest enrichment (~ 15.8 -fold) and accounts for 41% ($SE = 5.0\%$) of the heritability of general risk tolerance; the “H3K9ac peaks” category shows a 5.5-fold enrichment and accounts for 21% ($SE = 3.7\%$) of the heritability of general risk tolerance.

12.1.3 Results: Tissue types

The results of the tissue-level analyses are reported in **Extended Data Fig. 12.3a**, and **Supplementary Table 12.2**. For general risk tolerance, the enrichment of Central Nervous System is strongest (~ 2.94 -fold, $P = 5.90 \times 10^{-20}$), followed by Adrenal/Pancreas (~ 2.47 -fold, $P = 1.41 \times 10^{-7}$). The enrichment estimates for Immune/Hematopoietic, Cardiovascular, Liver, and Skeletal Muscle are also significant, and also survive Bonferroni correction. Our estimates imply that SNPs bearing the Central Nervous System and Adrenal/Pancreas annotations account for 43.8% ($SE = 2.8\%$) and 23.1% ($SE = 2.5\%$) of the heritability of general risk tolerance, respectively; the corresponding figure for Immune/Hematopoietic is 37.6% ($SE = 3.6\%$). These estimates are all highly significant and survive Bonferroni correction for ten tests.

However, the enrichment statistics are potentially misleading because of possible confounding. For example, many SNPs bear multiple annotations. It is thus of interest to examine the τ_c estimates of each tissue (i.e., the coefficients from the stratified LD Score regression). The τ_c for a given tissue type and phenotype corresponds to the effect of a one-unit increase in a SNP’s stratified tissue-specific LD score on the expected square of the SNP’s GWAS estimate from the phenotype’s GWAS (where the SNP genotype has been standardized), after controlling for the annotations from the baseline model. It is also an estimate of the expected increase in the phenotypic variance accounted for by a SNP due to the SNP’s being in the given tissue category, controlling for the annotations from the baseline model. SNPs that bear a tissue annotation with a large and positive τ_c will tend to account for a larger share of a phenotype’s heritability.

Extended Data Fig. 12.3a shows, for general risk tolerance and height, the ratio of the τ_c estimates over the LD Score estimates of phenotypic heritability^{III, mmm}. For each phenotype, we normalized the τ_c ’s and their standard errors by the LD Score heritabilityⁿⁿⁿ to increase comparability across phenotypes. Here, only Central Nervous System ($z = 8.81$, $P = 6.04 \times 10^{-19}$) and Immune/Hematopoietic ($z = 3.19$, $P = 1.42 \times 10^{-3}$) have positive coefficients surviving Bonferroni correction (note that these analyses excluded SNPs in the MHC region on chromosome 6). We also verified that the coefficient for Immune/Hematopoietic remained positive and highly significant ($z = 4.71$, $P = 2.47 \times 10^{-6}$) even after controlling for Central Nervous System (in addition to the baseline annotations) in the LD Score partitioned regression. Thus, the positive coefficient for Immune/Hematopoietic is not due to possible overlap between the Immune/Hematopoietic and Central Nervous System annotations.

In these respects, general risk tolerance differs notably from a physical trait such as height, for which Connective/Bone has the largest positive z -score ($z = 6.08$, $P = 1.20 \times 10^{-9}$) and Central

^{III} Based on the LD score framework, $h_{LDSC,y}^2 = \sum_c M_c \tau_{c,y}$, where y denotes the phenotype and M_c is the number of SNPs with annotation c among the SNPs used to calculate the LD scores. Thus, normalizing the τ_c ’s by $h_{LDSC,y}^2$ is equivalent to normalizing the τ_c ’s by a weighted sum of the τ_c ’s, where the weights are given by the number of SNPs with the different annotations.

^{mmm} The confidence intervals were obtained with the delta method, assuming zero covariance between the τ_c ’s and the phenotypes’ LD score heritabilities.

ⁿⁿⁿ Each phenotype’s LD score heritability was obtained from the phenotype’s LD score regression⁵³.

Nervous System is the only tissue-level annotation with a *negative* coefficient^{94,000}. Finally, we note that the coefficients for Cardiovascular, Gastrointestinal, Liver, and Skeletal Muscle (which were significantly enriched) have *negative* τ_c coefficients for general risk tolerance. This means that the GWAS signal is actually negatively enriched for histone marks in these tissues, after taking the baseline annotations into account.

12.1.4 Discussion

The results of our baseline annotation model are similar to those reported by Finucane *et al.*⁹⁴ for a set of nine phenotypes and to those reported by Okbay *et al.*^{16,18} for educational attainment, subjective well-being, depressive symptoms, and neuroticism. That is, evolutionary conserved regions (i.e. conserved in mammals¹⁹⁶) bear the most significant enrichment, followed by annotations relating to modification of gene expression. This is in line with a large body of literature that suggests that complex traits are likely to mainly be affected by genetic variants involved in the modification of gene expression, rather than by direct coding variants¹⁹⁷. In particular, the significant enrichment of “H3K9ac histone marks” points to the involvement of active gene promoters in general risk tolerance¹⁹⁸.

The evolutionarily conserved category denotes regions of the genome that have been conserved in mammals throughout evolution. The evolutionary conservation of a genomic region can be studied through assessment of the “mutation rate” of such a region. That is, some regions of the genome accumulate base-pair substitutions more slowly than predicted by a model of selective neutrality¹⁹⁶, which implies that mutations in such regions tend to have deleterious effects that decrease evolutionary fitness, and are therefore subject to natural selection.

As for the tissue enrichments, we obtained significant τ_c estimates for Central Nervous System and Immune/Hematopoietic cell types. As explained above, this implies that risk-tolerance associated SNPs that are in or near histone marks in these tissues contribute relatively more to the per-SNP heritability of risk tolerance than most other SNPs. Because risk tolerance is a psychological trait, central nervous system enrichment is not surprising. More interesting is the current lack of significance of the τ_c estimates for Adrenal/Pancreas. This suggests the highly significant enrichment estimate for Adrenal/Pancreas is due to overlap with SNPs bearing other annotations rather than to the partial effect of bearing the Adrenal/Pancreas annotation, and may possibly question the role for the stress-response system in risk tolerance^{199–201}. We note, however, that the lack of significance of the τ_c estimates for Adrenal/Pancreas could also be due to lack of statistical power or to imprecisions or imperfections in the SNP annotations. The stress-response system is driven by the hormones cortisol and (nor)epinephrine (also known by the name (nor)adrenaline), which are all produced in the adrenals.

To the best of our knowledge, we are the first to find significant LDSC tissue enrichment of the immune system in a psychological or neuropsychiatric trait (although enrichment of *specific* immune pathways and/or immune tissue enhancer regions has been implicated in schizophrenia^{58,202}, bipolar disorder²⁰³ and major depression²⁰⁴). We note that involvement of the immune system is not corroborated by our other analyses (aside from the large number of genes

⁰⁰⁰ We caution that a detailed quantitative comparison of the results for the height and the general-risk-tolerance GWAS might be misleading, because the sample size of the height GWAS is smaller than that of the risk-tolerance GWAS, and because the heritability of height is substantially higher than the heritability of general risk tolerance. Nonetheless, our results suggest that different tissue types tend to be relatively more enriched for the two phenotypes.

significant in the MHC region across out MAGMA, SMR and lookup analyses), and it is unclear how to interpret this finding.

12.2 Identifying genes associated with general risk tolerance with MAGMA

To identify genes that are significantly associated with general risk tolerance, we conducted a gene analysis with MAGMA¹⁸⁰ (defined in **Supplementary Information section 11.2**) for each of ~18,000 genes, in a hypothesis-free manner. We then used the Gene Network²⁰⁵ co-expression database to gain insight into the functions of the significant MAGMA genes.

12.2.1 Gene analysis with MAGMA: Testing ~18,000 protein-coding genes

The gene-based analysis was performed with MAGMA¹⁸⁰ using the summary statistics from our meta-analysis of the discovery and replication GWAS of general risk tolerance (after applying genomic control using the intercept of the LD Score regression).

All the SNPs from these summary statistics were annotated to genes based on NCBI 37.3.13 gene definitions. Only SNPs located between the transcription start and stop sites of a gene were annotated to that gene. A per-gene test statistic is calculated as the mean of the GWAS $-\log_{10}(P)$ values for all the SNPs between the transcription start and stop sites of a gene; MAGMA then calculates a P value for the resulting gene test statistic, using a procedure that takes into account the non-independence of the SNPs within the gene due to LD²⁰⁶. We used our main reference panel (described in **Supplementary Information section 2.4**) as reference data to estimate LD. Bonferroni correction was applied to account for multiple testing, counting each gene as an independent test.

The SNP-to-gene annotation yielded 18,224 protein-coding genes containing at least one SNP present in the current GWAS. After Bonferroni correction for 18,224 tests, 285 genes showed a significant association with general risk tolerance (**Supplementary Table 12.3**). We will henceforth refer to these as the “MAGMA genes.” The top ten MAGMA genes are: *CADM2* (chr. 3: 85 Mb, $P = 1.08 \times 10^{-50}$; all the P values reported in this paragraph are Bonferroni-corrected P values), *MSRA* (chr. 8: 9 Mb, $P = 1.61 \times 10^{-28}$), *XKR6* (chr. 8: 10 Mb, $P = 3.54 \times 10^{-23}$), *FOXP2* (chr.7: 113 Mb, $P = 9.94 \times 10^{-19}$), *MFHAS1* (chr. 8: 8 Mb, $P = 2.76 \times 10^{-18}$), *RPILI* (chr. 8: 10 Mb, $P = 2.04 \times 10^{-15}$), *FOXP1* (chr. 3: 71 Mb, $P = 2.23 \times 10^{-13}$), *RBFOX1* (chr. 16: 5 Mb, $P = 1.99 \times 10^{-12}$), *ARNTL* (chr. 11: 13 Mb, $P = 2.27 \times 10^{-11}$), and *ERII* (chr. 8: 9 Mb, $P = 7.49 \times 10^{-11}$).

The results show that several long-range LD or inversion regions of the genome contain a disproportionate number of MAGMA genes (which is not surprising, since MAGMA does not correct for LD between genes in its gene-based test). For instance, the Human Leukocyte Antigen (HLA) region (and its surrounding area) carries a relatively large number of significant genes: there are 29 significant genes in the chr.6: 25-32 Mb region, of which 11 are zinc finger genes, six are histone protein genes, and three are olfactory function genes (none of the *HLA*-genes themselves are significant, however). The large candidate inversion region on chromosome 8 (chr.8: 8 to 12 Mb, previously associated with neuroticism and depressive symptoms¹⁸) and its surrounding area carry 21 significant genes, 13 of which are among the top 30 most significant MAGMA genes, and of which *MSRA* is the most significant by far (Bonferroni-corrected $P = 1.61 \times 10^{-28}$). A long-range LD region on chromosome five (chr.5: 135-138 Mb) contains 10 significant

genes, of which *ETFL* is the most significant by far (Bonferroni-corrected $P = 7.46 \times 10^{-7}$). Finally, we note that several of these genes (*MSRA*, *ERII*, *XKR6*, and *ETFL*) were also significant in the SMR analyses, although only in the analysis of blood *cis*-eQTLs, and not in the analysis with GTEx data on *cis*-eQTLs from brain regions (lack of replication with the brain eQTL data may be because these data are based on relatively small samples and statistical power was consequently limited).

12.2.2 Using co-expression to predict gene function

We used a co-expression database called Gene Network²⁰⁵ (accessed 6 September 2017) to gain further insight into the functions of the MAGMA genes. The advantage of Gene Network lies in its ability to assign functions to a gene, even if that gene has not been annotated with a function in an actual empirical experiment (as described below). This is an important advantage because many human genes have not yet been studied in depth and hence have poorly understood functions. Therefore, looking up only the genes with empirically established functions in databases is in most cases not fully informative.

The gene functions used by Gene Network are derived from expert-curated sets of genes known as “gene sets” or “pathways.” The experts involved in the curation of such gene sets continuously review the empirical literature of a gene, and assign a gene its functions based on empirical results from laboratory experiments. Gene Network then assigns genes “membership” to these expert-curated gene sets by establishing “guilt-by-association” based on their co-expression with genes that are known members of such gene sets. A total of 14,461 functionally defined gene sets served as input, taken from standard databases such as Gene Ontology²⁰⁷ (GO), Reactome^{208,209}, and Kyoto Encyclopedia of Genes and Genomes²¹⁰ (KEGG). These co-expression patterns also serve as input for DEPICT¹⁹² (**Supplementary Information section 12.4**). All members of the functionally defined gene sets or pathways share a particular biological function, which can range from very specific and bottom-up functions (e.g., “clathrin binding”) to more coarsely defined and top-down functions (e.g., “hormone activity”). The biological function can also refer to the cellular site where the gene performs its function (e.g., “neuromuscular junction”).

We briefly describe the Gene Network method here, noting that in-depth documentation of the method can be found in Fehrmann *et al.*²⁰⁵ and Pers *et al.*¹⁹². The Supplementary Information of Okbay *et al.*¹⁸ also contains a detailed description. Gene Network uses information from a total of 77,840 publicly available human, mouse, and rat Affymetrix microarrays from the Gene Expression Omnibus. These microarrays measure expression levels of at least 19,997 human genes (or, in the case of mouse and rat experiments, their mouse- or rat-orthologs). Gene co-expression levels were measured in Pearson correlation coefficients. The resulting correlation matrix was broken down into principal components (PCs), which gave rise to a total of 2,206 reliable “transcriptional components” (TCs).

Each PC (i.e. transcriptional component) captures a shared pattern of gene expression across experiments, implying shared biology of the genes that load highly on the PC. The PCs can capture subtle patterns of co-expression that potentially reflect shared biological pathways among genes, which would typically be overshadowed by strong global transcriptomic effects, and thus missed in a straightforward correlation analysis.

Subsequently, the authors of this method tested whether the pattern of co-expression captured by the PC significantly differentiated members of the gene set from all other genes, by testing the

difference between the mean PC loading of the gene set versus the mean loading of all genes *not* in the gene set, using a Welch *t*-test for unequal variances. Each row of the resultant $14,461 \times 2,206$ matrix of *t*-statistics (each element corresponding to a gene set and PC) was then correlated with the PC loadings of the individual genes, to test whether the expression patterns of each gene and gene set were correlated. Finally, the thresholds for declaring a correlation between a gene's PC loadings and a gene set's *t*-statistics as statistically significant were chosen to satisfy False Discovery Rate (FDR) < 0.05. Thus, each gene's membership to each gene set was tested, allowing significant associations to be queried from the Gene Network database.

From this database, we first recorded for each MAGMA gene all statistically significant results (i.e., all gene sets that are significantly correlated with the genes' PC loadings), derived from gene sets and pathways from the Gene Ontology²⁰⁷ (<http://amigo.geneontology.org/amigo>) domains Biological Process, Cellular Compartment, and Molecular Function. We also recorded the results listed under the Reactome²⁰⁸ (<http://www.reactome.org/>) pathways and under the Kyoto Encyclopedia of Genes and Genomes²¹⁰ (KEGG, <http://www.genome.jp/kegg/>) pathways. In each case, we report (in **Supplementary Table 12.4**) the five most frequently occurring gene sets across the MAGMA genes.

Gene Network also reports organs, tissues, and cell types in which the gene under investigation is significantly and specifically expressed (compared to other tissues). For each MAGMA gene, we recorded all these organs, tissues and cell types where the area under the receiver operating characteristic curve (AUC), with respect to the discriminating power of measured gene expression, exceeded 0.80 according to Gene Network. Gene Network derived this AUC from the difference between the samples of the focal tissue and all other tissues in the distribution of the queried gene's expression level. These differences were determined by text-mining the descriptions provided by experimenters who uploaded the expression data to the Gene Expression Omnibus (GEO). Note that the tissue/cell type labels taken from the Medical Subject Headings (MeSH) database can refer to different levels of a hierarchy and are therefore not necessarily mutually exclusive.

We note that the method has three potential inherent drawbacks. First, only genes with reliable gene expression data (i.e., which have survived quality control) are included in the database. Therefore, the co-expression matrix might exclude genes that may be important but that have not yielded reliable expression data. For instance, *FTO* (a gene implicated in obesity) is a notable gene not included in the database.

Second, the method does not assign functions to genes that have unique patterns of gene expression but that may well play a role in known biological processes.

The third potential drawback of our querying method is that some of the significant MAGMA genes may only capture signal from other, nearby MAGMA genes, and not have any independent causal effects of their own. In this analysis, we conduct the search for each of the MAGMA genes and sum their predicted gene functions as if they were independent. This potential dependence among genes must, however, be taken into account when interpreting the predicted gene function counts.

For each of the three Gene Ontology (GO) domains, the Reactome pathways, the KEGG pathways, and the significant organs, tissues, and cell types, **Supplementary Table 12.4** lists the five most frequently occurring predicted gene functions, cellular components, and cell-type/tissues for the 266 MAGMA genes that were present in the Gene Network database. The most frequently

occurring terms overwhelmingly point to neural function and anatomy. To a lesser degree, functions related to gene expression are also highlighted.

The top five tissues are all brain tissues: prefrontal cortex, which is predicted for 88 genes, frontal lobe (81 genes), visual cortex (81 genes), parietal lobe (80 genes), and putamen (which is part of the basal ganglia; 80 genes). The most interesting of these tissues is likely the prefrontal cortex, which is crucial for inhibition of emotion and planning of behavior²¹¹, and is one of the last brain regions to mature during pubertal development²¹².

The top terms for GO biological process and GO molecular function are “glutamate signaling pathway” (12 genes) and “glutamate receptor activity” (15 genes), respectively, while the top occurring terms for GO cellular component refer to dendritic (22 genes) and synaptic (22 genes) cell components. GO biological process also infers several functions related to modification of gene expression, namely “chromatin modification and organization” (11 genes) and “histone modification” (9 genes). The top terms for Reactome are “neuronal system” (20 genes), with the remainder of top terms referring to synaptic and neurotransmitter functions. Four out of five terms for KEGG are neural, with “calcium signaling pathway” (20 genes) as the top term. The only non-neural annotation for KEGG is “spliceosome” (18 genes).

12.3 Identifying genes associated with general risk tolerance with SMR

12.3.1 Background

The rationale for the Summary-based Mendelian Randomization (SMR) test²¹³, which is one of several methods that test for an association between gene expression and trait variation (e.g., refs.^{213–215}), is that GWAS associations for complex traits and common diseases are enriched in non-coding regulatory regions of the genome (e.g., ref.⁵⁸), suggesting that many causal variants influence trait variation through regulation of gene expression. There is also evidence that causal variants often do not target the gene closest to the association signal (e.g., ref.²¹⁶).

SMR differs from our eQTL lookups of top GWAS SNPs (**Supplementary Information section 12.6.3**) in several important ways. First, it differs in its starting point: instead of testing *all top GWAS SNPs* for association with a probe, it tests *all gene expression probes* with a significant SNP association (i.e., all significant *cis*-eQTLs in a certain tissue dataset) for association with the phenotype. To this end, it uses SNPs as instrumental variables for gene expression levels to estimate the association between gene expression levels and the phenotype (as described in further detail below).

In this sense, SMR is more powerful than the eQTL lookups, where the number of tests is informed by the stringent GWAS *P* value threshold (as only genome-wide significant SNPs are put forward for eQTL testing). In SMR, the number of tests is informed by the number of significant eQTLs, which runs in the thousands. In this way, SMR can in principle uncover many more gene-phenotype associations than the eQTL lookups.

In addition, the “HEIDI” test feature of SMR is able to discard eQTL associations that are caused by mere linkage, thereby discarding genes that are “spuriously” associated with the phenotype (as described in further detail below). This feature distinguishes SMR from other transcriptome-wide analysis methods, and from the eQTL lookups, which may uncover genes that are merely associated with the phenotype due to linkage. We still perform the eQTL lookups, however, as they can give us some insight into the functional annotation of our GWAS top hits.

12.3.2 Methods

Summary statistic-based Mendelian Randomization (SMR)²¹³ is a method for estimating the effect of an exposure (x), such as gene expression, on a phenotype of interest (y), using summary-level data on estimated SNP effects (z , the instrumental variable) from large-scale GWAS and eQTL studies. In a Mendelian Randomization (MR) framework, the effect of x on y (b_{xy}) is:

$$\hat{b}_{xy} = \hat{b}_{zy} / \hat{\beta}_{zx},$$

where \hat{b}_{zy} is the estimated SNP effect from a GWAS for trait y and $\hat{\beta}_{zx}$ is the estimated SNP effect on the expression level of gene x from an independent eQTL study.

The sampling variance of \hat{b}_{xy} is given by:

$$\text{var}(\hat{b}_{xy}) \approx \frac{b_{zy}^2}{\beta_{zx}^2} \left[\frac{\text{var}(\hat{\beta}_{zx})}{\beta_{zx}^2} + \frac{\text{var}(\hat{b}_{zy})}{b_{zy}^2} - \frac{2\text{cov}(\hat{\beta}_{zx}, \hat{b}_{zy})}{\beta_{zx} b_{zy}} \right],$$

where $\text{cov}(\hat{\beta}_{zx}, \hat{b}_{zy})$ is assumed to be zero if β_{zx} and b_{zy} are estimated in independent cohorts. If $\hat{\beta}_{zx}$ and \hat{b}_{zy} are used in place of β_{zx} and b_{zy} (which are unknown), then the SMR test statistic, which is approximately χ^2 , takes the form:

$$T_{SMR} = \hat{b}_{xy}^2 / \text{var}(\hat{b}_{xy}) \approx \frac{z_{zy}^2 z_{zx}^2}{z_{zy}^2 + z_{zx}^2},$$

where z_{zy} and z_{zx} are the estimated z statistics from the GWAS and eQTL studies, respectively.

A significant SMR test result could be due to causality (i.e., if the effect of z on y is via x), pleiotropy (i.e., if z has pleiotropic effects on x and y) or linkage (i.e., if z_1 affects x and z_2 affects y , and z_1 and z_2 are in linkage disequilibrium). Like all MR methods, SMR is unable to distinguish between causality and pleiotropy on the basis of a single genetic instrumental variable (i.e., the top *cis*-eQTL as opposed to multiple *cis* and *trans*-eQTLs), and so we do not distinguish between pleiotropy and causality in our analyses.

On the other hand, it is possible, under some assumptions, to distinguish pleiotropy (or causality) from linkage using the HEIDI (Heterogeneity In Dependent Instruments) test developed by Zhu *et al.*²¹³. The rationale for the HEIDI test is that if the same causal variant has a pleiotropic effect on gene expression and the trait, then the estimate of b_{xy} should be identical for any SNP in LD with the causal variant. Therefore, a test for linkage is equivalent to testing for heterogeneity in the estimates of b_{xy} for the top *cis*-eQTL and any other significant SNPs in the *cis*-eQTL region. If there are m such variants (excluding the top *cis*-eQTL), and if we define $d_i = b_{xy(i)} - b_{xy(top)}$ and $\hat{\mathbf{d}} = \{\hat{d}_1, \hat{d}_2, \dots, \hat{d}_m\}$, then the HEIDI test against the null hypothesis $\mathbf{d} = 0$ takes the form:

$$T_{HEIDI} = \sum_i^m z_{d(i)}^2,$$

where $z_{d(i)}$ equals $\hat{d}_i / \sqrt{\text{var}(\hat{d}_i)}$. SNPs with $r^2 > 0.9$ with the top *cis*-eQTL are excluded from the set of m SNPs in the HEIDI test, because statistical power to distinguish between linkage and pleiotropy is limited if the causal variants have close to perfect LD. The test also excludes SNPs with eQTL P values $> 1.6 \times 10^{-3}$, due to the need to remove weak instrumental variables.

For a complete description of the SMR and HEIDI tests, see Zhu *et al.*²¹³.

12.3.3 Data

SMR requires summary statistics from large-scale GWAS and eQTL studies and genotype data from an ancestry-matched reference sample for estimating linkage disequilibrium. We used estimates of SNP effects (b_{zy}) from the meta-analysis of our discovery and replication GWAS of general risk tolerance, and we used summary data on *cis*-eQTLs (b_{zx}) from the eQTLgen Consortium's meta-analysis of 14,115 whole blood and peripheral blood samples (in prep.). We excluded probes in the MHC (26.1 to 33.8Mb on chromosome 6 based on hg19) due to the complexity of linkage disequilibrium in this genomic region, and we excluded SNPs with MAF < 0.01 and INFO score < 0.3. Analyses were restricted to probes with at least one *cis*-eQTL (defined as +/-1Mb from the middle of the probe) with $P < 5 \times 10^{-8}$, because Mendelian randomization assumes that the instrumental variable has a strong effect on the exposure. After filtering we had access to summary data on 12,751 probes and 10,209,777 (1000 Genomes imputed) SNPs. We used Bonferroni correction to account for multiple testing, resulting in a genome-wide significance threshold for the SMR test of $P < 3.9 \times 10^{-6}$ ($= 0.05/12,751$).

We based our primary analyses on eQTLs identified in blood (in the eQTLgen data) because the available sample size is at least an order of magnitude larger than for any other human tissue, and so this strategy maximizes statistical power for genes with shared genetic control of regulation across tissues. However, genetic control of regulation of some genes is tissue-specific, and so for each of the 21 probes passing both the SMR and HEIDI tests in the eQTLgen analysis, we also performed SMR analyses using estimates of SNP effects on gene expression in the human brain. We used summary eQTL data from the Genotype Tissue Expression Consortium (GTEx) for 11 brain regions (anterior cingulate cortex, caudate basal ganglia, cerebellar hemisphere, cerebellum, cortex, frontal cortex, hippocampus, hypothalamus, nucleus accumbens basal ganglia, pituitary, putamen basal ganglia) with sample size exceeding 70 (GTEx release v6p). We mapped Illumina expression probes in eQTLgen to GTEx transcripts using Ensembl gene IDs. We applied a relaxed eQTL P value threshold for inclusion of transcripts in the analysis of brain eQTLs ($P < 2.5 \times 10^{-3}$), due to the small number of transcripts tested per tissue, and we used a correspondingly lower significance threshold for the SMR test ($P < 2.16 \times 10^{-4}$, based on correction for testing of 21 probes in each of 11 brain regions).

We used our main reference panel (**Supplementary Information section 2.4**) for LD estimation. All analyses used hg19 coordinates.

12.3.4 Results and discussion

We used SMR to test for association between gene expression in blood and general risk tolerance using summary data from our GWAS meta-analysis and blood *cis*-eQTLs from the eQTLgen consortium. We identified 42 genes (tagged by 46 probes) at experiment-wide significance ($P < 3.9 \times 10^{-6}$), of which 19 (tagged by 21 probes) passed the HEIDI test, indicating we could not reject the null hypothesis of a single causal variant with pleiotropic effects on both risk tolerance and gene expression in blood (**Supplementary Table 12.5**). Thirteen of the 19 significant genes (68%) were in reported genome-wide significant loci for general risk tolerance, whereas the remaining six (*CTNNA1*, *SIL1*, *ZCCHC7*, *HNRNPK*, *HINFP*, *LYZ*) were “novel” discoveries, inasmuch as no reported general-risk-tolerance GWAS locus was located within 0.5Mb. These

latter discoveries point to genetic loci that are likely to be identified with genome-wide significance in larger GWAS of general risk tolerance. Several genes passing the SMR and HEIDI tests, including the top ranked gene *MSRA* ($P_{\text{SMR}} = 1.8 \times 10^{-13}$, $P_{\text{HEIDI}} = 8.8 \times 10^{-2}$), *ERII* ($P_{\text{SMR}} = 1.1 \times 10^{-7}$, $P_{\text{HEIDI}} = 8.5 \times 10^{-2}$) and *XKR6* ($P_{\text{SMR}} = 4.3 \times 10^{-10}$, $P_{\text{HEIDI}} = 0.59$), were also significantly associated with general risk tolerance in the MAGMA analysis (**Supplementary Information section 12.2**).

For each of the 21 probes passing both the SMR and HEIDI tests in the eQTLgen blood analysis, we repeated the SMR analysis for general risk tolerance using summary data on *cis*-eQTLs from the 11 GTEx v6p brain regions (**Supplementary Table 12.5**). Three genes passed both the SMR and HEIDI tests in one or more brain regions: *CTNNA1* (chromosome 5, 138.11Mb) was significant in caudate basal ganglia, cerebellum and cerebellar hemisphere; *CENPV* (chromosome 17, 16.25Mb) was significant in putamen basal ganglia, nucleus accumbens basal ganglia, caudate basal ganglia, anterior cingulate cortex (BA24), hippocampus and hypothalamus; and *ZSWIM7* (chromosome 17, 15.89Mb) was significant in cerebellum, cerebellar hemisphere, cortex, anterior cingulate cortex (BA24) and frontal cortex (BA9). These brain regions overlap with those identified by the DEPICT tissue prioritization analysis (see **Supplementary Information section 12.4**). For *CTNNA1* and *CENPV* (**Extended Data Fig. 12.2**), the risk-tolerance increasing allele was associated with decreased gene expression in blood and each of the respective brain regions. For *ZSWIM7*, the risk-tolerance increasing allele was associated with increased gene expression in blood, cerebellum and cerebellar hemisphere, but the results were inconsistent for cortex, anterior cingulate cortex and frontal cortex (i.e., the risk-tolerance increasing allele was associated with decreased expression). These findings suggest that *CTNNA1* and *CENPV* are putatively functional genes for risk-taking behavior but that more investigation is required to confirm the association with *ZSWIM7*.

According to the GTEx portal website, *CENPV* is highly expressed in many tissues, but especially in the brain's cerebellum, while *CTNNA1* is not particularly strongly expressed in brain tissue (compared to the other tissues). The function of *CENPV* has not been studied frequently, but two studies have discovered functions in cell division²¹⁷ and migration²¹⁸. *CTNNA1*, on the other hand, has been studied widely for its role in tumor suppression (e.g. ref.²¹⁹), and mutations in this gene are known to lead to retinal pigment dystrophy, causing macular disease²²⁰. Both *CTNNA1* and *CENPV* were significantly associated in the MAGMA gene based test, but neither were prioritized by DEPICT at FDR > 0.20. Therefore, the putative roles *CENPV* and *CTNNA1* may play in general risk tolerance remain unclear.

12.4 Prioritization of tissues, gene sets, and genes using DEPICT

12.4.1 Background and methods

DEPICT (Data-driven Expression Prioritized Integration for Complex Traits) is a computational tool that uses a set of GWAS summary statistics as input and allows enrichment analysis of tissues and gene sets and prioritization of genes in the associated loci (see ref.¹⁹² for details).

For this work, we used DEPICT release 194^{PPP}. DEPICT was run using the summary statistics of the meta-analysis of the discovery and replication GWAS of general risk tolerance (after applying

^{PPP} DEPICT release 194 handles GWAS data imputed with 1000 Genomes Project reference data and was downloaded in February 2016 from:

genomic control using the intercept of the LD Score regression). Only SNPs with GWAS P values less than 10^{-5} were used as input, and DEPICT-defined loci were defined by clumping these SNPs (PLINK clumping parameters: `--clump-p1 1e-5 --clump-r2 0.1 --clump-kb 500`, using 1000 Genomes Project pilot phase data as reference). Locus boundaries were then defined using a LD r^2 threshold of 0.5, and overlapping loci were merged, yielding 464 autosomal loci comprising 1,060 genes. DEPICT was run using default settings: 500 permutations for bias adjustment; 50 replications for false discovery rate estimation; normalized expression data from 77,840 Affymetrix microarrays for gene set reconstitution; 10,968 reconstituted gene sets for gene set enrichment analysis; and testing 209 tissue/cell types assembled from Medical Subject Headings (MeSH) annotations from 37,427 Affymetrix U133 Plus 2.0 Array samples for enrichment in tissue/cell type expression^{205,192}. Furthermore, gene expression data from 37 RNA-sequenced tissues from the Genotype-Tissue Expression (GTEx) Consortium⁷⁶ were used for tissue enrichment analysis.

12.4.2 Results: Prioritized tissues, gene sets, and genes

We begin by reporting the results of the DEPICT tissue enrichment analysis, which we conducted using GTEx RNA-sequencing gene expression data as well as using microarray data from 209 tissues^{192,205}. The DEPICT tissue enrichment analysis identifies tissues in which genes near general-risk-tolerance-associated SNPs are significantly overexpressed relative to genes in random sets of loci matched by gene density.

Using GTEx RNA-sequencing gene expression data, we identified eight significantly enriched tissues (FDR < 0.01), which all represent different areas of the brain (**Fig. 3b** and **Supplementary Table 12.6**). The top three prioritized tissues were cortical brain regions, namely “Frontal Cortex (BA9),” “Anterior Cingulate Cortex (BA24)” and “Cortex.” The cortical regions BA24 and BA9 are respectively located in the ventral anterior cingulate area and in the dorsolateral and medial prefrontal cortex^{221,222}. These two regions had initially been respectively associated with memory and emotion processing, though more recent work stresses their role in executive functions such as “executive control of behavior,” “inferential reasoning,” and “learning and decision making”^{223–227}. The DEPICT tissue prioritization analysis with the GTEx data also points to subcortical regions, including notably the nucleus accumbens, caudate, and putamen, which form the basal ganglia²²⁸, as well as the amygdala. The basal ganglia and amygdala are known for their respective role in motor control (and its impairment in Parkinson’s disease), and in the processing of negative emotions, such as fear. However, recent evidence also supports a role for the basal ganglia and the amygdala in reward processing and decision-making^{229–232}.

Using microarray data from 209 tissues, we replicated the brain enrichment and identified overall 19 significantly enriched tissues (FDR < 0.01) which are all part of the central nervous system (CNS) (**Extended Data Fig. 12.3b** and **Supplementary Table 12.7**). Note that we also observed an enrichment for the retina, which is part of the CNS but in DEPICT is categorized as “Sense Organs.” This analysis additionally showed expression in deeper brain structures, such as the rhombencephalon and metencephalon. The metencephalon, also known as the midbrain, is the major location of dopamine neurons²³³; we note, however, that this exact location, known as substantia nigra, was not prioritized by DEPICT in the analysis of GTEx tissues (FDR > 0.20). Moreover, the results seem to have little specificity toward the reward and decision-making

https://data.broadinstitute.org/mpg/depict/depict_download/bundles/DEPICT_v1_rel194.tar.gz.

system, as they also show expression in other CNS regions with diverse functions (e.g., the visual cortex).

Next, we report the results of the DEPICT gene set enrichment and gene prioritization analyses. DEPICT “reconstituted gene sets” are predefined gene sets taken from several bioinformatic databases (including Gene Ontology, Reactome, and Kyoto Encyclopedia of Genes and Genomes) that have been reconstituted using co-expression data to represent the weight of evidence of a given gene belonging to the gene set—as opposed to the all-or-nothing representation for the predefined gene sets (see Pers *et al.*, 2015¹⁹², for details). Each gene’s membership score with respect to a given reconstituted gene set is simply the *z*-statistic of the correlation between the gene’s vector of transcriptional components (TC) loadings and the binary gene set’s *t*-statistics with respect to the TCs—the same *z*-statistics used for testing the statistical significance of the Gene Network predictions. We identified 93 reconstituted gene sets that are significantly enriched (FDR < 0.01) for genes found among the general-risk-tolerance-associated loci (**Supplementary Table 12.8**). We used the Affinity Propagation tool²³⁴ to cluster related reconstituted gene sets into a network diagram (**Fig. 3a**)^{qqq}. The most strongly enriched gene sets highlight pathways related to the CNS. These include gene sets for regulation of neuronal projections (most notably synaptic and dendritic development) and synaptic transmission. Several of the most significant pathways relate to synaptic transmission activity, in particular GABA and glutamate neurotransmitter signaling. We were able to prioritize at least one gene at 106 of 464 loci defined by DEPICT (23% of the loci) at FDR < 0.01. In total DEPICT prioritized 122 genes (**Supplementary Table 12.9**) across those 106 loci.

The most significantly enriched reconstituted gene set is “dendrite development.” The presence of dendrites distinguishes neurons from all other cell types; these unique structures bestow the ability to receive signals of high spatiotemporal resolution. We mention just one aspect of dendrite development implicated by our results: the coordination or mutual influence required to juxtapose the axonic and dendritic sides of the developing synapse. In certain hippocampal pyramidal neurons, recognition molecules of the NGL/NTNG family produce a compartmentalization of the dendritic tree by ensuring the innervation of distinct portions of the tree by axons from correspondingly distinct sources. *NTGN1* and *NTNG2* encode ligands for receptors encoded by *LRRC4C* (also called *NGL1*) and *LRRC4* (also called *NGL2*) respectively (*NTGN1* is not prioritized by DEPICT, but *NTNG2* and *LRRC4* are prioritized at FDR < 0.01 and *LRRC4C* is prioritized at FDR < 0.05). *LRRC4C* and *LRRC4* are members of the leucine-rich repeat family, whose overarching function may be the exploitation of both family size and alternative splicing to create unique molecular fingerprints enabling precision wiring between neurons^{235,236}. Incoming axons bearing *NTGN1* and *NTNG2* respectively somehow find their corresponding receptors (*LRRC4C*, *LRRC4*) and thereby innervate non-overlapping compartments on distal and proximal dendrites²³⁷. These leucine-rich repeats are synaptic cell adhesion molecules (CAMs, such as *CADM2*) that transmit signals to the dendritic interior upon trans-synaptic binding, and thus their compartmental distribution may bestow varying information-processing properties along the length of a dendritic tree.

Our results implicate a number of different neurotransmitter receptors embedded in the membrane of the mature dendrite. Glutamate is the most abundantly employed excitatory neurotransmitter in the brain. When glutamate released from the axon of a transmitting presynaptic neuron binds to a

^{qqq} The script to produce the network diagram can be downloaded from <https://github.com/perslab/DEPICT>.

fast-acting AMPA-type glutamate receptor embedded in the dendrite of the receiving postsynaptic neuron, the channel coupled to the receptor momentarily opens by a lock-and-key mechanism and permits a massive influx of Na^+ ions. This is the primary mechanism leading to rapid depolarization of the receiving neuron, which can lead in turn to the neuron firing its own signal along its axon. Our DEPICT-prioritized genes include *GRI1A1*, which encodes the most common subunit appearing in the AMPA-type glutamate receptor.

Two genes encoding subunits of NMDA-type glutamate receptors are among our DEPICT-prioritized genes (*GRIN2A*, *GRIN2B*). NMDA-type glutamate receptors act like coincidence detectors: they only open when glutamate has landed on the receptor and the postsynaptic membrane potential has been sufficiently depolarized by the opening of other excitatory channels to remove the Mg^{2+} block. The emerging picture is of a receptor whose task is not to respond immediately to any given input impinging on the dendrite but rather to admit an agent of change into the dendrite in response to a temporal coincidence of synapse-specific input and membrane depolarization.

So far, we have been discussing excitatory neurotransmission—that is, a signal from one neuron to the next that pushes the receiving neuron closer to its own firing threshold. Now we turn to inhibitory neurotransmission, which has the effect of pushing the receiving neuron further away from its firing threshold. The most important source of inhibition in the cortex is mediated by receptors for the neurotransmitter GABA. They produce inhibition in many cases by generating hyperpolarizing changes in membrane potential that counteract the depolarizing changes caused by Na^+ influx through opened glutamate receptors. GABA_A receptors mediate the effects of alcohol intoxication and drugs such as benzodiazepines and barbiturates, which have anxiolytic effects.

A GABA_A neuron typically establishes outgoing connections only with neurons in its local vicinity. A glutamatergic pyramidal neuron, in contrast, often sends its axon to neurons that are far away (e.g., the opposite hemisphere). When this local connectivity of GABA_A neurons is emphasized, they are called interneurons. Because it is not possible to construct an information-processing device from only inhibitory neurons, we can reasonably say that they play some supporting or modulatory role.

GABA_A receptor subunits are grouped into subfamilies known respectively as α , β , γ , δ , ϵ , θ , ϕ , and ρ ²³⁸, the first three of which are represented in our DEPICT-prioritized genes (*GABRA2*, *GABRA4*, *GABRB1*, *GABRG3*). A complete receptor is composed of five subunits; the combinatorial number of possible pentamers is quite impressive, even if we put aside alternative splicing. Some subunits of GABA_A receptors contain regulatory sites for phosphorylation and domains that interact with proteins such as the product of *GPHN* involved in receptor trafficking and localization at synapses²³⁹.

The importance of both excitatory and inhibitory neurotransmission is affirmed by the significant enrichment of gene sets such as “glutamate receptor activity,” “abnormal miniature excitatory postsynaptic currents,” and “ GABA_A receptor activity.” Note that the cluster named after the latter gene set contains the second-most significant gene set in the entire analysis (**Fig. 3a**).

Four out of the top five prioritized genes encode proteins with transcriptional regulatory activity, some known to play a role in brain development. The strongest prioritized gene, *FOXG1*, encodes a transcription factor known to cause the congenital variant of Rett-syndrome²⁴⁰, a severe neurodevelopmental disease. The second strongest prioritized gene, *ASH1L*, encodes a histone

lysine N-methyltransferase involved in chromatin modification and has been implicated in autism and neurodevelopmental disorders²⁴¹. The third strongest prioritized gene, *FAM190A* (alias *CCSER1*), is a relatively unknown gene with no previous reported associations to brain-related phenotypes. The fourth and fifth strongest prioritized genes, *FOXO6* and *NEUROD*, both encode transcription factors and have been implicated in Alzheimer's Disease and regulation of memory consolidation, respectively^{242,243}.

12.5 Competitive gene-set analysis with MAGMA: Testing gene sets related to GABA and glutamate neurotransmitters

12.5.1 Background

Since both the DEPICT pathway analysis and the Gene Network analysis of MAGMA genes both pointed to glutamate and GABA neurotransmitters, we decided to also perform an *ex post* MAGMA competitive gene-set analysis of relevant glutamate and GABA gene sets. This allowed us to directly compare the enrichment of these gene sets to that of the gene sets associated with dopamine, serotonin, testosterone, estrogen, and cortisol, which we also tested in a MAGMA gene set analysis described, as described in **Supplementary Information section 11.2**.

Similar to what we did in **Supplementary Information section 11.2**, we looked up all relevant glutamate and GABA gene sets in MSigDB v6.1^{trr} and merged the resulting sets into one glutamate and one GABA superset. In addition, we tested the GABA and glutamate gene sets which were significant ($FDR < 0.05$) in DEPICT (see **Supplementary Information section 12.4**), although we updated the genes in those sets according to the gene sets in MSigDB v6.1.

Thus, in total, we conducted a MAGMA gene set analysis to test four glutamate and GABA gene sets (Panel **B** in **Supplementary Table 11.2**):

- 1) All GABA sets significant in DEPICT ($FDR < 0.05$, six gene sets) or with more than 10 counts in the Gene Network analysis of MAGMA genes (one gene set), containing 68 unique genes
- 2) All glutamate sets significant in DEPICT ($FDR < 0.05$, 12 gene sets) or with more than 10 counts in the Gene Network analysis of MAGMA genes (zero gene sets), containing 159 unique genes
- 3) All relevant GABA sets in MSigDB v6.1 (13 gene sets), containing 134 unique genes
- 4) All relevant glutamate sets in MSigDB v6.1 (27 gene sets), containing 276 unique genes

We tested all four sets together, and MAGMA computed the family-wise error correction accordingly.

12.5.2 Results and discussion

None of the four gene sets were significant (lower Bonferroni-corrected $P = 0.299$, Panel **B** in **Supplementary Table 11.3**). We are unsure why this might be the case but, as we further discuss this in **Supplementary Information section 12.7.4**, our MAGMA competitive gene-set analysis

^{trr} See http://software.broadinstitute.org/cancer/software/gsea/wiki/index.php/MSigDB_v6.1_Release_Notes for details on this release

of the four glutamate and GABA gene sets has some likely limitations. First, the statistical power of a MAGMA gene set analysis fails to increase substantially with increasing sample sizes¹⁸⁷. Second, important effects of single SNPs in a gene set may be overshadowed by the high average P values of other SNPs in the gene or gene set. Third, it might be that only specific glutamate and/or GABA genes or pathways might be relevant for general risk tolerance, and that our merging of those pathways into major glutamate and GABA sets decreased our statistical power for discovery of relevant effects. (In the MAGMA gene-based analysis, only 12 of the 354 unique GABA/glutamate genes included in the four gene sets tested here were significant at Bonferroni-corrected $P < 0.05$.)

12.6 Lookup of protein-altering variants and *cis*-eQTLs

In this section, we conducted several analyses that annotate the 132 lead SNPs from the meta-analysis of the discovery and replication GWAS of general risk tolerance, to gain insights about the biological systems they may affect. (For consistency with the rest of the bioannotation analyses, we performed these analyses with the 132 lead SNPs of the meta-analysis combining the discovery and replication GWAS of general risk tolerance, and not with the 124 lead SNPs from the discovery GWAS.^{sss})

Specifically, we examined the overlap between these lead SNPs and various SNPs and genes that have been identified or annotated in previous GWAS and functional genomic databases. First, we checked whether our lead SNPs (and SNPs in LD with them) contain protein-altering variants. Second, we looked up the lead SNPs (and SNPs in LD with them) in an expression quantitative trait loci (eQTL) database.

12.6.1 SNPs in LD with lead SNPs

Since the genome is characterized by widespread linkage disequilibrium (LD), it is important to examine the overlap with SNPs that are in LD with our lead SNPs (as opposed to only considering our lead SNPs). Moreover, some data sources do not contain some of our lead SNPs but contain SNPs in strong LD with them, and a lead SNP does not actually have to be the causal SNP in the region: it may just be the most often genotyped or most accurately imputed SNP (i.e., the SNP measured with the least measurement error and therefore yielding a lower P value); it may tag an unmeasured causal SNP in the LD block; or it may simply be the most significant SNP in the block

^{sss} Because the lead SNPs of the discovery GWAS are not a perfect subset of the lead SNPs of the meta-analysis of the discovery and replication GWAS, we investigated the overlap of the identified lead SNPs. The discovery GWAS identified 124 lead SNPs, and the meta-analysis of the discovery and replication GWAS identified 132 lead SNPs; 97 of these are lead SNPs in both. We investigated if the non-overlapping lead SNPs of each GWAS are in weak LD with a lead SNP of the other (defined here as $r^2 > 0.1$). The discovery GWAS identified seven lead SNPs that are not in weak LD with a lead SNP in the meta-analysis of the discovery and replication GWAS, and the meta-analysis of the discovery and replication GWAS identified 15 lead SNPs that are not in weak LD with a lead SNP of the discovery GWAS. (The list of these seven and 15 SNPs is available upon request.) These seven and 15 lead SNPs are all highly significant ($P < 5E-7$) in the other GWAS. In conclusion, approximately 117 lead SNPs overlap across the two GWAS, and the 15 additionally identified lead SNPs in the meta-analysis of the discovery and replication GWAS are all highly significant in the discovery GWAS. The slightly larger number of lead SNPs in the meta-analysis of the discovery and replication GWAS was to be expected because of the increased statistical power due to the inclusion of the replication GWAS.

due to sampling variation. For these reasons, it is crucial to also consider the LD partners of the lead SNP. We obtained a list of SNPs in strong LD with our lead SNPs using the PLINK $-r2$ command in our main reference panel (described in **Supplementary Information section 2.4**). Here we used an LD cutoff of $r^2 > 0.6$ and a distance cutoff of 250kb from the lead SNP. We refer to these SNPs as the “LD partners” of our lead SNPs. In total, the clumping procedure designated 132 “lead SNPs,” which tagged a total of 8,199 LD partners.

12.6.2 LD with protein-altering variants

Using the Haploreg v4.1 database²⁴⁴ (download date 18 May 2016, <http://www.broadinstitute.org/mammals/haploreg/haploreg.php>), we ascertained the protein-altering status of our lead SNPs and their LD partners. More precisely, we ascertained whether our lead SNPs and their LD partners are annotated as “missense,” “nonsense,” “splice site donor/acceptor,” or “frameshift” in the 1000 Genomes Phase 3 European population. The Haploreg database uses annotations from dbSNP⁷³ to determine functional status. We examined these annotations because protein-altering variants have a higher *a priori* probability of being causally relevant: they can result in stronger phenotypic differences between individuals compared to non-protein-altering variants. Most of the genome is non-coding, which underlines the importance of the detection of protein-altering variants.

The results can be found in **Supplementary Table 12.10**. We found that there were eight protein-altering SNPs among the lead SNPs’ LD partners, including two of the lead SNPs themselves. All protein-altering variants were relatively common (with $0.11 < MAF < 0.50$) missense variants (meaning that they result in an amino acid change). The two missense lead SNPs are in genes *WSCD2* (on chromosome 12) and *NF1* (on chromosome 17). Little is known about *WSCD2*, but it is highly expressed in the brain and has predicted functions in long term memory and calcium signaling, according to Gene Network²⁰⁵. *NF1* is involved in the genetic disorder “neurofibromatosis,” which is characterized by widespread fibromatous tumor formation and cognitive disability²⁴⁵. Other missense variants in LD with our lead SNPs were found in *FREMI* and *BHLHE22*. *FREMI* encodes a protein that may play a role in craniofacial and renal development, while *BHLHE22* is thought to be involved in the differentiation of sensory neurons. The remaining four missense variants in LD with our lead SNPs were in the HLA-region, of which three were in the olfactory receptor gene *OR2J2*, and one in histone gene *HIST1H1T*.

12.6.3 eQTL lookup

The publicly available results from the Genotype-tissue expression project (GTEx, www.gtexportal.org, version v6p)⁷⁶ allow us to look up significant associations between our lead SNP and their LD partners and *cis*-gene expression in distinct human tissues. The GTEx team obtained postmortem samples from human donors and used RNA sequencing to measure RNA levels in distinct tissues from those samples. The GTEx team also genotyped the donors with SNP microarrays. The lookup of the risk-tolerance lead SNPs and their LD partners was performed on the publicly accessible summary statistics (downloaded 13 December 2016, version v6p) of the significant gene-*cis*-SNP pairs from GTEx. We only considered the set of significant *cis*-eQTLs (i.e., gene-*cis*-SNP pairs), where the *cis*-window is defined as a 1MB window around the gene’s transcription start site. In addition, when a gene transcript was a significant eQTL for more than one of our query SNPs, we only report the eQTL statistics (i.e., effect size, *P* value) for the SNP in highest LD with the lead SNP (which might be the lead SNP itself). That way, we only report

each significant eQTL gene once in each tissue (even though the gene may have appeared several times for the tissue, for several lead SNPs and LD partners). In cases where more than one lead SNP was an eQTL for the same gene, we report the eQTL for only one of them.

To identify statistically significant *cis*-eQTLs, the GTEx consortium computes a gene-specific nominal *P* value threshold for each gene, with a permutation method as described in more detail in ref.²⁴⁶. The eQTLs we report here are all statistically significant, in the sense that the nominal *P* value of the slope of the regression of the gene's expression value on the SNP was smaller than the gene-specific nominal *P* value threshold. This regression also included several technical covariates, such as the top principal components of the genetic relatedness matrix.^{ttt}

We only recorded findings for the following pre-selected tissues, which we suspected could be of biological relevance for risk tolerance: (1) all 11 brain tissues^{uuu} available in the GTEx data; (2) tibial nerve tissue; (3) thyroid tissue; (4) gonadal (i.e. testis and ovary) tissue; and (5) adrenal gland tissue. In addition, we queried (6) the “whole blood” tissue for eQTLs, as this tissue had the highest sample size. Since expression profiles in whole blood and the brain are moderately correlated, we might uncover potentially interesting eQTLs by using information from whole blood^{247,248}.

The donor samples sizes for the brain tissues range from $n = 72$ to 103, and sample sizes for the other five tissues range from $n = 126$ to 338. We included tibial nerve tissue (a peripheral nerve located in the lower leg) because the eQTL results for this tissue are based on a relatively large sample size (compared to the brain tissues) of $n = 256$. Since central and peripheral nervous tissues are anatomically similar, their expression profiles are likely to be comparable. Thyroid tissue is included because thyroid hormone is essential for healthy brain development, and might be associated with mood and cognition^{249,250}. Testis and ovary tissue are included, as these organs produce sex hormones (e.g., testosterone, estrogen) that might be involved in risk tolerance, as detailed in our literature review (**Supplementary Information section 11.1**). Adrenal tissue is included because the adrenals produce hormones associated with the general stress response (i.e., (nor)epinephrine and cortisol), which might be related to risk tolerance, as detailed in our literature review (**Supplementary Information section 11.1**). Note that the presence of eQTLs in these tissues *does not* imply statistical enrichment of the GWAS signal in these tissues; it merely shows that our lead SNPs are associated with gene expression in these tissues (which might simply be a result of pleiotropy, and which thus might not be of causal relevance to general risk tolerance).

Supplementary Table 12.11 reports the results. We find eQTL overlaps in every tissue (including all individual brain tissues) we queried. In total, we find 188 unique gene transcripts for 59 different lead SNPs. 82 of these gene transcripts only had Ensembl gene IDs and no gene symbols (i.e. letter symbol gene names such as “*HLA*” which is an abbreviation of Human Leukocyte Antigen, according to the HUGO gene nomenclature committee), most likely because they do not have named gene products (i.e., proteins). Out of the 188 unique gene transcripts, 47 lie in or near the human MHC (i.e., Major Histocompatibility Complex, also known as Human Leukocyte Antigen (HLA)) region (~chr. 6: 26-32 Mb), of which five are *HLA*-genes.

Finally, we note that there was one eQTL for a GABA gene (*GABRG1*, or GABA_A receptor gamma1 subunit), in which the general-risk-tolerance-increasing allele decreased expression of

^{ttt} For additional details, see <http://www.gtportal.org/home/documentationPage#staticTextAnalysisMethods> and ref.²⁴⁶.

^{uuu} These include pituitary tissue (which GTEx Portal does not list among brain tissues).

GABRG1 in tibial nerve tissue. We found no eQTLs for glutamate genes in any of the queried tissues.

12.7 Summary of main findings from the biological annotation analyses

The overall goal of the analyses described in this section was to gain insight into the biology of general risk tolerance. We employed two general approaches. First, we leveraged data at the genome-wide level to prioritize potentially relevant tissues and genomic annotations (using LDSC partitioning of heritability) and genes (using MAGMA gene-based analysis, SMR transcriptome-wide analysis, and DEPICT gene prioritization analyses). Second, we focused on the loci around the general-risk-tolerance lead SNPs (in the lookups of protein-altering variants and GTEx eQTLs) to gain insight into the biological functions of our lead SNPs. Here, we summarize the results of those analyses.

12.7.1 *Enrichment of SNPs annotated to the central nervous system*

First, we tested enrichment of the GWAS signal in 10 different tissues/cell-types using LDSC partitioning of heritability. After controlling for tissue overlap and correcting for multiple testing, we found significant positive involvement for the central nervous system and the immune system. Involvement of the central nervous system was expected (since general risk tolerance is a behavioral trait), but enrichment of immune cell types might be surprising. To the best of our knowledge, immune enrichment in LDSC partitioning analyses has not previously been reported for a behavioral or psychiatric trait, although we note again that enrichment of *specific* immune pathways and/or immune tissue enhancer regions has been implicated in schizophrenia^{58,202}, bipolar disorder²⁰³ and major depression²⁰⁴).

We note that enrichment of immune tissues or function did not come forward in any of the other analyses. Further research is therefore needed to corroborate a role for immune processes in general risk tolerance. Finally, we note that several additional tissues, such as liver and adrenal/pancreatic tissues, showed statistically significant enrichment, but that our estimates of their partial effects on enrichment (i.e., their τ_c coefficients) were not significant, thereby suggesting that their enrichment may have been primarily driven by other overlapping annotations. We therefore cannot claim a role for the adrenal system in general risk tolerance. However, this non-finding might be a result of the combination of adrenal with pancreas tissue, and the possible non-specificity of histone marks to these particular tissues. More fine-grained analyses are therefore needed to corroborate or refute a role for the adrenal system in modulating risk tolerance.

12.7.2 *Enrichment of SNPs near genes that are highly expressed in the frontal and prefrontal cortex*

While the LDSC partitioning results pointed to the central nervous system (which technically includes the brain, brainstem and spinal cord), results from the other analyses allowed us to gain deeper insight into which particular brain regions might be of relevance to general risk tolerance. Although the overlap between the genes prioritized by DEPICT and the significant MAGMA genes was not very high (only 71 out of 285 genes of the MAGMA genes were among the 215 genes prioritized at FDR < 0.05 by the DEPICT gene prioritization analysis), both the Gene Network functional analysis of the MAGMA genes and the DEPICT tissue enrichment analysis with the GTEx tissue-specific gene-expression data separately point to enrichment of the frontal

cortex, and in particular the prefrontal cortex. The frontal cortex is a large and functionally and anatomically complex brain area; it is the largest neocortical region in humans and primates, and receives and integrates neuronal input from both cortical and subcortical regions²¹¹. The posterior dorsal area of the frontal cortex contains the motor cortex, which is responsible for planning and directing bodily movement; damage to this area can result in perturbed motor function or paralysis. The prefrontal cortex (the most anterior part of the brain), on the other hand, is involved in planning, directing, and inhibiting emotion, behavior and cognition (broadly known as “executive function”). Damage to the prefrontal cortex has been well-documented due to the widespread practice of prefrontal lobotomy (a relatively simple surgical procedure) in 20th century America and with textbook patient cases of accidental lesion such as Phineas Gage. Damage to the prefrontal cortex can result in severe impairment marked by behavioral disinhibition and emotional deregulation, cognitive disability, impaired decision making, a tendency to display context-inappropriate behavior, and inability to plan future action²¹¹. The frontal cortex is also one of the last brain areas to mature during development (with full maturation reached around age 25)²¹², which has been postulated to underlie the social and behavioral disinhibition and increased display of risk-taking behaviors that characterizes puberty and adolescence^{251–253}.

At a finer resolution, the DEPICT tissue enrichment analyses with the GTEx tissue-specific gene-expression data show that genes near general-risk-tolerance-associated SNPs are highly expressed in the prefrontal cortex (BA24 and B9) and the basal ganglia (nucleus accumbens, caudate, putamen), compared to other organs and tissues. The DEPICT tissue enrichment analysis using the microarray data confirmed these CNS regions’ enrichment, and additionally showed expression in deeper brain structures like the midbrain, among other tissues. The Gene Network tissue enrichment analyses also point to enrichment of the prefrontal cortex, and of the putamen, which is part of the basal ganglia. Finally, two of the three genes that passed both the SMR and HEIDI tests, *CTNNA1* and *CENPV*, were significant in parts of the basal ganglia (*CTNNA1* was significant in the caudate and *CENPV* was significant in the putamen, nucleus accumbens, and caudate).

12.7.3 *A possible convergence with results from the neuroscientific literature*

Interestingly, the prefrontal cortex, the basal ganglia and the midbrain are the major components of the *cortical-basal ganglia circuit*, which includes and connects the reward system, the motor system, and the cognitive and behavioral control systems. This whole network of brain regions has been shown to be critically involved in learning, motivation, behavioral control, and decision-making, notably under risk and uncertainty, in both humans and non-human primates^{233,254}. Decision making under risk requires several cognitive processes (e.g. such as reward processing, fear, or higher executive control) underpinned by different neural systems which are coherently connected and integrated in current models of the *cortical-basal ganglia circuit*. In this complex circuit, the processing of reward and decision-making is presumed to be largely underpinned by dopamine^{230,255}. Dopaminergic neurons are mostly located in the midbrain (substantia nigra and ventral tegmental area) where they project to the basal ganglia and the prefrontal cortex, and appear to encode signals about past and future rewards, used to guide behavior^{230,255}.

There is therefore a remarkable concordance between the enriched CNS regions highlighted by the DEPICT analyses and the circuit in charge of reward processing, behavioral control, and decision-making defined by decades of behavioral and physiological studies in human and non-human primates²³³. It is however difficult to claim that this circuit is *specifically* tagged by the DEPICT

tissue enrichment analysis with the GTEx data, given that the GTEx data only cover a fraction of all brain regions, mostly restricted to brain regions overlapping with the cortical-basal ganglia circuit. Although these results, in line with a large body of neuroscientific studies, suggest a role of the cortical-basal ganglia circuit in the risk-tolerance phenotype, their inferential value should therefore be considered with caution.

12.7.4 *A likely role for glutamate and GABA neurotransmitters in the modulation of risk tolerance*

In addition to neuroanatomy, much speculation has revolved around the role of specific hormones and neurotransmitter systems in modulating general risk tolerance. Our review of the literature attempting to link risk tolerance to biological pathways (**Supplementary Information section 11**) identified five main biological pathways that have been tested by this literature: the steroid hormone cortisol, the monoamines dopamine and serotonin, and the steroid sex hormones estrogen and testosterone. Consistent with the lack of enrichment reported in **Supplementary Information section 11** for genes associated with these five pathways, neither the Gene Network analysis of the MAGMA genes nor the DEPICT gene set prioritization analysis point to these pathways. (As mentioned above, however, we note that our results are consistent with a role of the cortical-basal ganglia circuit in modulating risk tolerance, and that dopamine is presumed to play an important role in that circuit.)

On the other hand, both the Gene Network functional analysis of the MAGMA genes and the DEPICT gene set prioritization analysis point to a role for glutamate and GABA neurotransmitters, which are the main excitatory and inhibitory neurotransmitters in the brain, respectively²⁵⁶. They are colloquially referred to as the “workhorses” of the brain, as they are the brain’s principal neurotransmitters. At the time of writing, no other published large-scale GWAS of cognition, personality, or psychiatric phenotypes has pointed to clear roles for *both* glutamate and GABA, although we note that glutamate neurotransmission has been implicated in recent GWAS of schizophrenia—by recent analyses of common SNPs⁵⁸ and rare and *de novo* mutations^{257,258}—and of major depression²⁰⁴. That our results, unlike those of other well-powered GWAS, point to both of these neurotransmitters suggests that the relative balance between excitatory and inhibitory neurotransmission may be a relatively strong contributor to variation in risk tolerance across individuals.

We note that GABA-glutamate balance may be implicated in epilepsy, which we do not consider a cognitive, personality, or psychiatric trait. A certain class of GABAergic drugs (anticonvulsants) are used in the treatment of epilepsy, and GABA-glutamate balance has been postulated to underlie epilepsy pathophysiology²⁵⁹. In the largest epilepsy GWAS conducted to date²⁶⁰, two highlighted genes involved in GABA regulation (*GABRA2* and *PCDH7*, which contain or are near SNPs that reached suggestive or genome-wide significance in the epilepsy GWAS) were also highly significant in our MAGMA gene-based analysis (**Supplementary Information section 2612.2.1** and **Supplementary Table 12.9**). However, in our GWAS Catalog lookup, we found no overlaps between our general-risk-tolerance lead SNPs, and SNPs in LD with them, and any epilepsy lead loci^{vvv}.

^{vvv} We did find one overlap for one epilepsy SNP and an LD partner for one of the number of sexual partners lead SNPs (**Supplementary Information table 262.12**).

We note that none of the four glutamate and GABA gene sets we tested in an *ex post* MAGMA competitive gene-set analysis were significant (**Supplementary Information section 12.5**). We are unsure why this might be the case, but we note that this analysis had some likely limitations. First, its statistical power barely increases with increasing sample sizes¹⁸⁷. Second, a MAGMA competitive gene-set analysis might not be the optimal approach for discovery of subtle but phenotypically important genetic effects driven by regulatory regions, as this analysis effectively calculates a weighted average of GWAS $-\log_{10}(P)$ values across each gene in the gene set (where the average is weighted to correct for LD between SNPs). Hence, important regulatory effects of single SNPs may be overshadowed by the high average P values of other SNPs in the gene sets. A third possible explanation is that only specific glutamate and/or GABA genes or pathways might be relevant in general risk tolerance, and that our merging of those pathways into major glutamate and GABA supersets decreased our statistical power for discovery of relevant effects. (In the MAGMA gene-based analysis, only 12 of the 354 unique GABA/glutamate genes included in the four gene sets tested here were significant at Bonferroni-corrected $P < 0.05$.) Hence, the lack of significance of the four glutamate and GABA gene sets we tested in the *ex post* MAGMA competitive gene-set analysis does not nullify the evidence from our Gene Network and the DEPICT analyses that glutamatergic and GABAergic neurotransmission contributes to variation in risk tolerance across individuals.

12.7.5 *A possible link between the immune system and risk tolerance*

We now briefly discuss the LDSC immune tissue enrichment. We noted that, to the best of our knowledge, general risk tolerance is the first behavioral trait to show immune tissue enrichment in LDSC partitioned analysis according to tissue type, and that this enrichment was independent of SNPs in the MHC region. We also noted that none of our other analyses show immune enrichment—the Andersson²⁶¹ “FANTOM5” enhancers (which are indicative of immune activity⁹⁴) were not a significant genomic annotation in the LDSC partitioned analyses according to genomic annotation, and the DEPICT and Gene Network analyses of MAGMA genes also did not show immune pathways (although it is noteworthy that these bioannotation analyses excluded the MHC region). However, we do find GWAS hits in the MHC region, and several genes in this region came up in the MAGMA and GTEx lookup analyses. That is, 29 of the 285 genes significant in MAGMA and 47 of the 188 GTEx eQTL lookups were in (or around the borders of) the MHC region (while SMR excluded SNPs in the MHC region). While a role for the immune system in neuropsychiatric traits and related behavior is feasible given previous enrichment of immune pathways for psychiatric traits^{58,203,204,258}, we are unsure how to interpret a role for the immune system in general risk tolerance, given that our results do not unequivocally point in this direction. Researchers have previously theorized a role for a “behavioral immune system”^{262,263} in human behavior. This is seen to be complementary to the “real” immune system—that is, humans are said to employ risk averse behaviors to minimize their risk of infectious disease. An extension of this theory could be that individuals who are genetically prone to “weaker” immune systems might be more risk averse than others, but this is extremely speculative. Before accepting such conclusions, we encourage future bioannotation work to uncover which specific immune pathways might be involved in general risk tolerance.

12.7.6 The *CADM2* gene and risk tolerance

Finally, we briefly discuss the top locus from our discovery GWAS of general risk tolerance. The top lead SNP that marks this locus is located in the intronic region of *CADM2*, and is our most significant general-risk-tolerance lead SNP by far, with a P value that is fifteen orders of magnitude smaller than the second lead SNP's P value ($P = 2.1 \times 10^{-40}$ vs. $P = 7.6 \times 10^{-25}$). The *CADM2* locus was previously identified by a study using the relatively small sample size of the first release of UK Biobank data⁶⁶, and all of our six supplementary GWAS phenotypes identified lead SNPs within or near *CADM2* (**Supplementary Information section 3**). *CADM2* was by far the most significant MAGMA gene and was also prioritized by DEPICT at $FDR < 0.01$. However, we do note that only *CADM2*'s antisense RNA partner *CADM2-AS1* was a significant cis-eQTL gene in our GTEx lookup tissues, while *CADM2* failed the HEIDI-test for heterogeneity in the SMR analysis. This suggests that *CADM2* may not be the causal gene in the locus, or that its effect on general risk tolerance is not mediated by *CADM2* expression. However, we do stress that the brain eQTL and SMR analyses are strongly underpowered due to the relatively small samples sizes for brain tissues, and we can thus not properly study *CADM2* expression in relevant brain tissues.

According to GTEx, *CADM2* is overexpressed in the brain, and in particular in the frontal cortex. *CADM2* is a large gene (spanning more than 1 Mb) located in a long-range LD region, and has been associated with a myriad of behavioral phenotypes in previous GWAS (**Supplementary Information section 3**). *CADM2* encodes a member of the synaptic cell adhesion molecule family. Molecules in this family have been found to be crucial for synapse formation⁷⁴ and plasticity⁷⁵, and have been associated with autism²⁶⁴ and impaired social behavior²⁶⁵. However, *CADM2* itself has mainly been studied for its potential role in tumor progression²⁶⁶; its specific role in the brain is unclear. We therefore propose *CADM2* as an interesting candidate gene for future follow up work.

13 Additional information

13.1 Author contributions

Jonathan Beauchamp, Philipp Koellinger, and Daniel Benjamin designed and oversaw the study. Jonathan Beauchamp closely supervised the analyses and led the writing of the manuscript.

Richard Karlsson Linnér was the lead analyst, responsible for quality control, the meta-analyses, summarizing the overlap across the results of the various GWAS, and the SNP heritability analyses. Edward Kong conducted the population stratification, replication, and proxy-phenotype analyses.

Robbee Wedow led the genetic correlation analyses, and Mark Fontana contributed to those analyses. Pietro Biroli led the polygenic score prediction analyses, and Richard Karlsson Linnér, Edward Kong, Robbee Wedow, Abdel Abdellaoui, Ronald de Vlaming, Mark Fontana, and Michel G Nivard, contributed to those analyses. Pietro Biroli and Christian Zünd conducted the MTAG analyses.

The bioinformatics analyses were led by Fleur Meddens. Analysts who assisted Fleur include: Jacob Gratten and Maciej Trzaskowski (SMR), Anke Hammerschlag (MAGMA), Gerardus Meddens (Gene Network) and Pascal Timshel (DEPICT). Maël Lebreton conducted the review of the literature attempting to link risk tolerance to biological pathways.

Richard Karlsson Linnér prepared the majority of figures; Edward Kong and Stephen Tino also prepared some figures. Aysu Okbay, Cornelius A Rietveld, and Stephen Tino helped with several additional analyses.

David Cesarini, Jacob Gratten, and James Lee provided helpful advice and feedback on various aspects of the study design.

All authors contributed to and critically reviewed the manuscript. Pietro Biroli, Richard Karlsson Linnér, Edward Kong, Fleur Meddens, and Robbee Wedow made especially major contributions to the writing and editing.

13.2 Cohort-level contributions

Cohort	Author	Study design & mgmt.	Data collection	Geno-typing	Geno-type prep.	Pheno-type prep.	Data analysis
23andMe	Pierre Fontanillas						X
23andMe	Aaron Kleinman						X
23andMe	Adam Auton	X					
23andMe	David A. Hinds	X					
Add Health	Jason D Boardman		X				
Add Health	Kathleen Mullan Harris		X				
Add Health	Matthew B McQueen		X				
Army STARRS	Murray B Stein	X	X				
Army STARRS	Chia-Yen Chen			X	X		X
Army STARRS	Sonia Jain					X	
Army STARRS	Robert J Ursano	X	X				
BASE-II	Lars Bertram	X	X	X	X		
BASE-II	Christina M Lill		X	X			X
BASE-II	Peter Eibich		X			X	X
BASE-II	Gert G Wagner		X			X	
HRS	Ronald de Vlaming					X	X
NFBC1966	Ville Karhunen					X	X
NFBC1966	Andrew Conlin	X	X			X	
NFBC1966	Minna Männikkö	X	X			X	
NFBC1966	Rauli Svento	X	X			X	
NTR	Michel G Nivard					X	X
NTR	Abdel Abdellaoui				X		X
NTR	Conor V Dolan	X					
NTR	Dorret I Boomsma	X	X		X		
RS	Frank J van Rooij		X			X	X
RS	Albert Hofman	X					
RS	Mohammad A Ikram	X	X				
RS	Andre G Uitterlinden			X	X		
RS	Henning Tiemeier	X	X			X	
RS	Patrick JF Groenen	X					X
RS	A Roy Thurik	X					X
RS	Cornelius A Rietveld				X	X	X
STR	Cornelius A Rietveld				X	X	X
STR	Patrik KE Magnusson	X	X	X	X		
STR	Robert Karlsson			X	X		
STR	Magnus Johannesson	X	X			X	
STR	David Cesarini	X	X			X	
UKB	Mark Alan Fontana				X	X	X

UKB	Richard Karlsson Linnér				X	X	X
UKB	Robbee Wedow					X	
UKB	Cornelius A Rietveld				X	X	X
UKHLS	Yanchun Bao				X		X
UKHLS	Melissa C Smart				X	X	
UKHLS	Jon White			X			
UKHLS	Meena Kumari	X					
Viking	Paul RHJ Timmers						X
Viking	Peter K Joshi						X
Viking	David W Clark					X	X
Viking	James F Wilson	X	X				
ZMI	Ernst Fehr	X	X			X	
ZMI	Daniel Schunk	X	X	X		X	
ZMI	Arcadi Navarro	X			X		
ZMI	Laura Buzdugan					X	X
ZMI	Urs Fischbacher		X				
ZMI	Gregor Hasler	X					
ZMI	Gerard Muntané				X		X
ZMI	Klaus M Schmidt		X				
ZMI	Matthias Sutter		X				
ZMI	Carlos Morcillo-Suarez						X

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