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Citation for published version:
Hernandez, DG, Nalls, MA, Moore, M, Chong, S, Dillman, A, Trabzuni, D, Gibbs, JR, Ryten, M, Arepalli, S, Weale, ME, Zonderman, AB, Troncoso, J, O'Brien, R, Walker, R, Smith, C, Bandinelli, S, Traynor, BJ, Hardy, J, Singleton, AB & Cookson, MR 2012, 'Integration of GWAS SNPs and tissue specific expression profiling reveal discrete eQTLs for human traits in blood and brain' Neurobiology of disease, vol 47, no. 1, pp. 20-28. DOI: 10.1016/j.nbd.2012.03.020

Digital Object Identifier (DOI):
10.1016/j.nbd.2012.03.020

Link:
Link to publication record in Edinburgh Research Explorer

Document Version:
Peer reviewed version

Published In:
Neurobiology of disease

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Integration of GWAS SNPs and tissue specific expression profiling reveal discrete eQTLs for human traits in blood and brain

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Abstract

Genome wide association studies have nominated many genetic variants for common human traits, including diseases, but in many cases the underlying biological reason for a trait association is unknown. Subsets of genetic polymorphisms show a statistical association with transcript expression levels, and have therefore been nominated as expression quantitative trait loci (eQTL). However, many tissue and cell types have specific gene expression patterns and so it is not clear how frequently eQTLs found in one tissue type will be replicated in others. In the present study we used two appropriately powered sample series to examine the genetic control of gene expression in blood and brain. We find that while many eQTLs associated with human traits are shared between these two tissues, there are also examples where blood and brain differ, either by restricted gene expression patterns in one tissue or because of differences in how genetic variants

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There is no conflict of interest to disclose.

Datasets

Datasets have been submitted to GEO: accession # GSE36192

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are associated with transcript levels. These observations suggest that design of eQTL mapping experiments should consider tissue of interest for the disease or other trait studied.

Introduction

Genome-wide association (GWA) studies have provided novel insights into human traits by identifying single nucleotide polymorphisms (SNPs) associated with disease, including type 1 diabetes, coronary artery disease, HIV-1 infection and type 2 diabetes (Fellay et al., 2007; Preuss et al., 2010; Scott et al., 2007; Sladek et al., 2007; Steinthorsdottir et al., 2007; Todd et al., 2007a; Yang et al., 2010; Zeggini et al., 2008), or other phenotypes. Because GWAS identify loci rather than functional variants, most GWAS have provided limited insights into underlying mechanisms (Hindorff et al., 2009). Therefore, annotating the possible functional effects of genetic risk variants is important in understanding genomic data.

Mapping of expression quantitative trait loci (eQTL) is one way to demonstrate that a risk variant within a locus has a functional effect on gene expression (Cheung et al., 2005; Morley et al., 2004; Myers et al., 2007; Stranger et al., 2007). eQTL analysis is performed by examining the association of each SNP with expression of mRNA transcripts. In general, eQTL effects are stronger for SNPs and transcripts that are physically close to each other (Gibbs et al., 2010). Trait associated SNPs from GWAS have been proposed to be more likely associated with expression differences than other SNPs (Nicolae et al., 2010). Such studies have generally been performed with transformed cell lines but eQTLs can also be identified in liver (Schadt et al., 2008), kidney (Wheeler et al., 2009), cell lines from asthma patients (Dixon et al., 2007; Moffatt et al., 2007) blood (Nalls et al., 2011a), subcutaneous adipose tissue (Emilsson et al., 2008) and brain (Gibbs et al., 2010; Heimzen et al., 2008; Liu et al., 2010a; Myers et al., 2007; Webster et al., 2009). For at least some loci, eQTLs are found consistently in both transformed cells and in primary tissues (Bullaughey K, 2009). Overall, this data might suggest that functional annotation of GWAS loci can be performed in any convenient tissue.

Studying brain tissue is particularly challenging because these tissue samples have to be collected post mortem and there is a high degree of cellular heterogeneity. Although some eQTLs have been nominated for brain diseases, such as MAPT in Parkinson’s disease (PD) and progressive supranuclear palsy (PSP) (Hoglinger et al., 2011; Nalls et al., 2011b; Tobin et al., 2008; Vandrovcova et al., 2010), many nominated loci for brain phenotypes are not functionally annotated.

To explore the tissue specificity of eQTLs, we analyzed expression in brain and blood using SNPs abstracted from the NHGRI catalog of GWAS. We specifically wanted to address whether it is necessary to examine brain tissue to detect eQTLs for brain traits, including neurological diseases and psychiatric events, or whether the same information could be obtained from a more accessible tissue such as blood. We find that while many eQTLs are shared between blood and brain, there are specific instances, not always simply related to tissue specific gene expression levels, where the tissue studied limits detection of eQTLs.

Material and Methods

Samples

Fresh, frozen tissue samples from the frontal lobe of the cerebral cortex and from the cerebellum were obtained from neurologically normal Caucasian subjects. Genomic DNA was extracted using phenol-chloroform and RNA using Trizol from subdissected samples (100–200mg). Peripheral blood specimens were collected using PAXgene tubes. RNA was
extracted from peripheral blood samples using the PAXgene Blood mRNA kit (Qiagen, Crawley, UK) according to the manufacturer’s instructions.

**Genotyping and Imputation**

Genotyping was performed using the Illumina Infinium HumanHap550 v3, Human610-Quad v1 or Human660W-Quad v1 Infinium Beadchip and common SNPs across all platforms were identified for each sample. SNPs were excluded if they showed < 95% genotyping success rate per SNP, minor allele frequency (MAF) < 0.01 or Hardy-Weinberg equilibrium (HWE) p-value < 1E-7. Quality control was carried out using PLINK v1.07 for each cohort separately prior to imputation and was determined by comparing the subjects reported gender with the genotypic gender determined using PLINK’s check sex algorithm.

Ethnicity and cryptic relatedness was determined using Identity-by-State (IBS) clustering and multidimensional scaling analyses within PLINK using genotypes that had been merged with data from HapMap Phase III, ASW, TSI, CEU, JPT, CHB and YRI populations [http://hapmap.ncbi.nlm.nih.gov/]. The subset of SNPs used were shared across studies, using only common SNPs that are not correlated within a 50 SNP sliding window at an $r^2 > 0.20$, with each window overlapping by 5 SNPs. Samples were clustered using multi-dimensional scaling, removing outliers > 3 standard deviations from the mean component vector estimates for C1 or C2 for the combined CEU and TSI samples. Cryptically related samples were excluded after pairwise identical by descent estimates were calculated, excluding any samples sharing greater than a 0.15 proportion of alleles.

Markov Chain based haplotyper (MACH 1.0.16) was used to impute non-assayed genotypes for blood and brain datasets independently using the June 2010 release of the 1000 Genomes Project build-36 reference panel, using default settings for MACH. Imputed SNPs were excluded from the analysis if their minor allele frequency (MAF) was <0.01 and if their $r^2$ was <0.3

**GWAS SNPs**

Trait and disease associated SNPs were extracted from the NHGRI catalog of published GWAS at http://www.genome.gov/gwastudies/ on July 30th 2011. Analyses was restricted to the following criteria: discovery p-value < 5E-08, initial sample size >1000 (or 1000 cases in binomial analyses), replication sample size >500 (or 500 cases in binomial analyses), number of SNPs >100,000, samples of European ancestry and risk allele frequency of SNP(s) greater or equal to 0.01.

**Expression Profiling**

Expression profiling was performed largely as previously described (Gibbs et al., 2010). RNA was biotinylated and amplified using the Illumina® TotalPrep-96 RNA Amplification Kit and directly hybridized onto HumanHT-12_v3 Expression BeadChips. Where possible, the same RNA samples were used from our previous study that used HumanRef8 Expression BeadChips. Raw intensity values for each probe were normalized using cubic spline in BeadStudio (Illumina) then log2 transformed. Individual probes were included in analysis if they were detected (P<0.01) in more than 95% of samples in the series.

To define probes within +/- 1MB of SNPs, probes were re-annotated using ReMOAT (http://www.compbio.group.cam.ac.uk/Resources/Annotation/). Ambiguous probes that mapped to multiple positions, or were identified as having design problems in ReMOAT, were excluded from subsequent analyses. To remove potential bias resulting from polymorphisms, all probes that included an analyzed SNP within the 50mer probe were removed.
Expression QTL Analyses

Starting with 447 subjects in the brain series, after data normalization and quality control, the brain mRNA dataset included 399 samples and ~9000 mRNA probes that were detected in >95% of all samples. The blood dataset started with 712 samples, of which 501 passed all our QC steps; 5094 mRNA probes were detected in >95% of samples. In each brain region, mRNA probes within 500kb of the chromosomal location of each SNP were incorporated into linear regression modeling using MACH2QTLv1.08. Estimates of the association between the allelic dose of each SNP as a predictor of proximal gene expression levels were generated. These linear regression models were adjusted for biological covariates of age at death and gender, the first 2 component vectors from multi-dimensional scaling, as well as methodological covariates including post mortem interval (PMI), tissue bank and hybridization batch. SNPs with fewer than 3 minor homozygotes detected (based on either genotyped SNPs or maximum likelihood genotypes from imputation) were excluded from analyses. A consensus set of results was extracted from the frontal cortex, cerebellum and blood eQTL datasets with identical overlapping combinations of GWAS SNPs and proximal cis mRNA probes. Significant associations were determined within each tissue type using a 5% FDR adjustment for multiple testing. Proportions of tested associations were calculated per tissue based on this subset of the eQTL results, and were compared using simple chi-squared tests.

Case studies of specific loci

Identical statistical models were utilized to test our ability to detect known-associated eQTLs in previously published reports in tissues not previously investigated in GWAS. Results for these loci were mined for all associations within each +/- 500kb region around top SNPs within each locus from the published GWAS within each tissue.

Results

Power to detect eQTLs in large blood or brain datasets

Directly comparing expression datasets derived from brain and whole blood in human samples is difficult because brain samples are taken post mortem whereas blood samples are routinely taken during life. Therefore, we used two large, well-powered series from different sets of individuals to maximize our ability to find eQTLs in each tissue type. For brain, we expanded our previous dataset (Gibbs et al., 2010) in frontal cortex and cerebellum and obtained whole blood from 712 individuals from the InCHIANTI study (Wood et al., 2011). For consistency, we used the same expression array platform (Illumina HT-12 beadchips containing 48,000 probes) for all samples. After quality control, the brain mRNA dataset included 399 samples with data at 9000 probes. The blood dataset included 501 samples containing expression data from 5094 probes. Following imputation and quality control, ~2.2 million SNPs were available for analysis in all sample sets.

Because the final number of samples within the blood and brain groups differed, we performed post-hoc power calculations to compare ability to detect eQTLs (Fig. 1). Based on our previous work in brain (Gibbs et al., 2010), the strength of the association varies substantially for different eQTLs. Therefore, we estimated power over a range of minor allele frequencies and of effect sizes for the eQTLs, using Z as a measure of effect size standard deviations of difference for each minor allele under an additive model. As an example of power in the two datasets at a realistic pair of these parameters, the blood dataset had 98.8% power to detect eQTLs at an effect allele frequency of 0.2 and an additive effect size of Z=0.5 whereas the brain dataset had 93.9% power to detect the same magnitude of effect. This analysis demonstrates that the difference in power in the two datasets is minimized as the fraction of true eQTL effect sizes rises. For eQTLs with moderate effect...
sizes (Z>0.2) we were reasonably powered in both series; therefore, we proceeded to compare the ability to detect eQTLs in both datasets.

**Gene expression in blood versus brain in human populations**

It is expected that gene expression profiles would be divergent between blood and brain tissues but similar for two brain regions. To test this, we ranked as percentiles the normalized gene expression values averaged for all subjects, setting non-detected probes to zero. Gene expression values were shown to be highly divergent between blood and either frontal cortex or cerebellum tissue for a large number of genes that were only detected reliably in one tissue or the other (Fig. 2A,B). In contrast, gene expression was more similar between frontal cortex and cerebellum and there were fewer uniquely expressed genes (Fig. 2C). Analysis using percentile ranked variance rather than mean values for each probe yielded similar results (Fig. 2D–F), showing that mean expression and variance in expression were closer in the two brain regions than in blood.

**eQTL discovery for genes expressed in blood and brain**

We next examined the relative ability of the three datasets to detect eQTLs from regions nominated in GWAS. We abstracted SNPs associated with human traits based on the NHGRI catalog of GWA, yielding 1366 loci. Of these, 783 SNPs passed the criteria of having a replicated association with traits or diseases and being within 0.5MB of the chromosomal position of a probe for gene expression. We chose the threshold of 0.5MB based on previous data (Gibbs et al., 2010) where we saw the average distance between a SNP and significant eQTL was 121Kb and >90% of significant eQTLs were detected within 0.5MB.

We manually annotated the traits studied in each GWAS as related to blood (176 SNPs), brain (61 SNPs) or other (546 SNPs) phenotypes (Supplementary file 1). For example, we annotated traits associated with neurological or psychiatric conditions as “brain” and markers of subtypes of blood cell markers as “blood”. We then used this list of SNPs to perform eQTL analysis. We first performed the eQTL analysis in a uniform way by only considering the subset of probes and SNPs detected in all tissue types, or 2929 SNP:probe pairs. This analysis identified eQTLs that were highly significant in all three tissues and additional eQTLs distinctly significant in either blood or brain tissues (Fig. 3). Of the shared eQTLs, three stood out as highly significant in all three tissues for three SNPs including a single mRNA probe, ILMN_1695585 that maps to the **RPS26** gene on chromosome 12q13.2, within 500KB of three GWAS SNPs associated with Type 1 diabetes (False discovery rate (FDR) corrected P<1.45×10^{-38} for association with rs11171739 in the frontal cortex, P<6.72×10^{-51} in cerebellum and P<9.46×10^{-67} in blood) (Barrett et al., 2009, 2007; Cooper et al., 2008; Hakonarson et al., 2008; Todd et al., 2007b). Additional significant SNP:probe pairs found in both datasets included SNPs associated with traits such as mean corpuscular volume (Ganesh et al., 2009), smoking behavior (2010), eye color (Liu et al., 2010b), plasma levels of liver enzymes(Yuan et al., 2008) and inflammatory bowel disease (Kugathasan et al., 2008) (Supplementary Table 1). For fifteen SNP:probe associations that were significant in brain and blood, the direction of effect was consistent across all three tissues.

A divergent set of eQTLs were found in the blood dataset when compared with cerebellum and frontal cortex (Fig. 3A,B). Several of these eQTLs were for probe ILMN_1666206, which maps to the **GSDML** gene on Chr17q12. These correlations are linked with five separate GWA studies associating Type 1 diabetes (2007; Barrett et al., 2009; Cooper et al., 2008; Hakonarson et al., 2008; Todd et al., 2007b), Crohn’s disease (Barrett et al., 2008) and Ulcerative colitis (McGovern et al., 2010) to the same locus (Supplementary Table 1).
Additionally, the ILMN_1666206 probe has been nominated as underlying an eQTL in studies of asthma and white blood cell traits associated with a pro-inflammatory state (Moffatt et al., 2007; Nalls et al., 2011a). The FDR corrected \( p \) value for the most significantly associated SNP, rs2290400, with this probe was \( 6.41 \times 10^{-32} \) in blood but 0.924 and 0.896 in the cerebellum and frontal cortex respectively.

To examine this phenomenon further, we compared all SNPs within the GSDML/ORMDL3 region to expression of ILMN_1666206 in all tissues. Although expression was detected, we did not find significant associations with any SNPs in the brain, but found strong associations between probe expression and proximal SNPs in blood tissue (Fig. 4B). This locus therefore represents an example of a blood-specific eQTL.

Conversely, a subset of SNP:probe pairs reached significance in the brain samples but not in blood (Fig. 3A,B and Supplementary Table 1). For example, rs713586, which was nominated for association with body mass index (PMID: 20935630), was significantly associated with expression of ILMN_1676893 in cerebellum (FDR corrected \( p = 6.09 \times 10^{-5} \)) and frontal cortex (FDR corrected \( p = 1.53 \times 10^{-5} \)) but showed no association in blood (FDR corrected \( p = 0.89 \)). This probe maps to the adenylate cyclase gene ADCY3 on chromosome 2 (Fig. 4A). Interestingly, variation in ADCY3 has been nominated in a number of GWAS including for alcohol dependence (Edenberg et al., 2010) and major depression (Wray et al., 2010).

Overall this data suggests that while some eQTLs are consistent between tissues, there is a subset where a genetic effect on gene expression exists in one tissue context but not the other, despite probe detection in both instances.

eQTL discovery for genes with expression restricted to blood or brain

Given that there were differences in gene expression between tissues (Fig. 2), we next analyzed eQTLs unique to each tissue by examining the association of GWAS SNPs with expression of probes detected in either blood or brain but not in both (Supplementary Table 2).

In blood, there was a highly significant (FDR corrected \( p = 1.27 \times 10^{-131} \)) association between rs2549794 at a Crohn’s disease locus (Franke et al., 2010) and expression of ILMN_1743145, which maps to the LRAP/ERAP2 gene (Fig. 5A). Other associations measurable only in the blood datasets include rs2304130 and ILMN_2134224, and rs6120849 and ILMN_2402805. These SNPs were nominated as associated with measurements of total cholesterol and protein C respectively in plasma (Tang et al., 2010; Waterworth et al., 2010).

We also found a series of significant associations in brain and not blood. Several of these associations were probes on chromosome 12 associated with rs11171739 or rs1701704, two SNPs nominated for Type-I diabetes. Additional associations included a series of SNPs on chromosome 17. Previous studies have noted an effect of chromosome17 SNPs on expression of the MAPT gene that is associated with risk of PD and PSP (2011; Hoglinger et al., 2011). We therefore examined expression of ILM_1710903, which maps within the coding sequence of MAPT, with association of SNPs across Chr17 and saw robust signals in both the frontal cortex and cerebellum (Fig. 5B). This effect was driven by the H1/H2 haplotype across the MAPT locus as conditioning the analysis on a proxy SNP decreased the apparent eQTL signal.
Collectively, these results show that while some eQTLs are shared across tissues, there are examples where restricted expression levels in one tissue limit the ability to detect significant associations.

**Overall ability to detect eQTLs depends on tissue type and gene expression**

We next compared the proportions of eQTLs found in the three sample series and considered whether these were associated with brain, blood or other phenotypes (Table 1). In the analysis restricted to probes detected in both blood and brain tissues, 125 eQTLs were found within 500 KB of any GWA SNP in blood, versus 40 eQTLs found in the brain dataset. Of these, 21 significant eQTLs were found in the blood dataset for blood traits, while 16 eQTLs were found in blood for brain traits. Six significant eQTLs were found in brain, counting either cerebellum or frontal cortex, for brain traits and an additional six eQTLs were found in brain for blood traits out of 40 total significant associations. The proportions of blood and brain traits with detected eQTLs were similar in blood and brain samples (two tailed Z-test, P=0.18, 1.0 respectively).

We performed independent analyses of probes that were only detected in one tissue as more probes were tested in brain (1877 in frontal cortex and 1853 in cerebellum) than in blood (413). There were 107 significant associations in cerebellum and 90 in frontal cortex compared to 21 in blood but we did not find over-representation of traits annotated as brain related in the brain datasets (Table 2).

**Discussion**

We have performed an eQTL analysis using SNPs from the NHGRI catalog of GWAS in two tissue types, blood and brain (frontal cortex and cerebellum). The nominated SNPs are associated with a variety of human traits, including diseases, physiological markers such as blood cell numbers and continuous traits such as height. We specifically addressed whether it is necessary to examine brain tissue to detect eQTLs for brain phenotypes or whether the same information could be obtained from blood. We find that while many eQTLs are shared between blood and brain tissues, there are specific instances, not always simply related to gene expression levels, where the detection of eQTLs is limited by the tissue studied.

A small number of eQTLs are detectable in all three datasets tested. A proportion of these common eQTLs demonstrated strong effects sizes, such as SNPs associated with Type 1 diabetes on chromosome 12 (Barrett et al., 2009; Burton et al., 2007; Cooper et al., 2008; Hakonarson et al., 2008; Todd et al., 2007b) or associated with smoking behavior on chromosome 19 (Furberg H, 2010). We have therefore demonstrated that coincident eQTLs exist between blood and brain tissues and therefore discrete eQTLs are found in more than one human primary tissue as previously suggested (Bullaughey K, 2009; Ding et al., 2010; Greenawalt et al., 2011).

There were eQTLs that could be detected only in one tissue type and in some cases these are due to differences in gene expression. This is true for genes such as MAPT, which encodes for the tau protein that is expressed largely in post-mitotic neurons. Therefore, there will be cases where, when interrogating GWAS data it will be important to examine the target tissue of interest, thus affirming the need to look at brain for studies related to neurological or psychiatric phenotypes.

Of greater interest is that we also found a subset of eQTLs that appear to be tissue specific, despite the probes being reliably detected in all samples series. It is possible that genetic variants can affect expression levels exclusively in a subset of tissues. For example, gene expression may be altered in a tissue- and timing-specific manner by cis- regulatory
elements (Cooper et al., 2008). In this case, although multiple tissues may be permissive for expression, different –cis regulatory elements are being employed in each tissue and lead to quantitatively different expression levels. Understanding why there are examples where differences in expression do not explain eQTL detection in a simple way will be an important question for future studies.

One caveat to these studies is that direct comparison of datasets derived from separate tissue types with differential ascertainment methods is difficult. Specifically, the brain samples were taken from deceased subjects whereas blood samples were drawn in life. However, post mortem interval has been shown not to be a major confound within brain expression data (Gibbs et al., 2010; Trabzuni et al., 2011) and we corrected for this and other known methodological variables in the statistical model. However, because the samples used here were from different individuals, we cannot exclude that we are detecting rare alleles and/or genetic variants on a background of common SNPs. As demonstrated by power analysis, the current dataset is not powered to directly detect rare alleles but has good power to detect relatively large eQTL effect sizes. Therefore, this analysis performs best for loci that are tagged by common variants and where the effect of the minor allele on expression is relatively large. It is also important to note that in the present study, we limited our analysis to transcripts within a relatively narrow (0.5MB) window around each SNP. This is larger than the average distance between SNP and associated transcript of 121Kb (Gibbs et al., 2010) but may inadvertently omit true eQTLs at larger distances while maintaining power. Larger series would be needed to expand the analyses to more distal effects.

Further dissection of such loci will likely require deep sequencing of the genome for many individuals and additional large-scale studies. One general limitation of hybridization based arrays is that detection of low expression genes is difficult, which may be overcome by RNA sequencing in the future. In addition to both of these technological developments, eQTL surveys such as the one presented here will need to be repeated as the numbers of SNPs nominated by GWAS studies increases. This is perhaps particularly true for brain related phenotypes. Although we did not find that there were significantly more eQTLs for brain phenotypes using brain expression data, the number of replicated GWAS ‘hits’ for neurological and psychiatric conditions is still quite small and we might expect brain to be more sensitive as the number of replicated loci increases. We have not tested all possible SNPs in the current analysis to maintain power to detect significant associations, but such analyses could be performed on an ad hoc basis for nominated SNPs in future GWAS without the loss of power caused by testing the whole genome.

Overall, we demonstrate a number of clear and key examples where brain tissue is required for eQTL discovery. We conclude that functional studies in one tissue have the capacity to inform our understanding of regulatory variation in general, but that there are sufficient numbers of counter-examples to suggest that for neurological and psychiatric traits we should continue to examine gene expression in the brain.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This research was supported in part by the Intramural Research Program of the NIH, National Institute on Aging (Z01-AG000947 and Z01-AG000185) and in part by the UK Medical Research Council. The InCHIANTI study baseline (1998–2000) was supported as a “targeted project” (ICS110.1/RF97.71) by the Italian Ministry of Health and in part by the U.S. National Institute on Aging (Contracts: 263 MD 9164 and 263 MD 821356).
References

Barrett JC, et al. Genome-wide association study and meta-analysis find that over 40 loci affect risk of type 1 diabetes. Nat Genet. 2009; 41:703–7. [PubMed: 19430480]

Barrett JC, et al. Genome-wide association defines more than 30 distinct susceptibility loci for Crohn’s disease. Nat Genet. 2008; 40:955–62. [PubMed: 18587394]

Bullaughey KCC, Coop G, Gilad Y. Expression quantitative trait loci detected in cell lines are often present in primary tissues. Hum Mol Genet. 2009; 18:4296–4303. [PubMed: 19671653]

Burton PR, et al. Genome-wide association study of 14,000 cases of seven common diseases and 3,000 shared controls. Nature. 2007; 447:661–78. [PubMed: 17554300]

Cheung VG, et al. Mapping determinants of human gene expression by regional and genome-wide association. Nature. 2005; 437:1365–9. [PubMed: 16251966]

Cooper JD, et al. Meta-analysis of genome-wide association study data identifies additional type 1 diabetes risk loci. Nat Genet. 2008; 40:1399–401. [PubMed: 18978792]

Ding J, et al. Gene expression in skin and lymphoblastoid cells: Refined statistical method reveals extensive overlap in cis-eQTL signals. Am J Hum Genet. 2010; 87:779–89. [PubMed: 21129726]

Dixon AL, et al. A genome-wide association study of global gene expression. Nat Genet. 2007; 39:1202–7. [PubMed: 17873877]

Edenberg HJ, et al. Genome-wide association study of alcohol dependence implicates a region on chromosome 11. Alcohol Clin Exp Res. 2010; 34:840–52. [PubMed: 20201924]

Emilsson V, et al. Genetics of gene expression and its effect on disease. Nature. 2008; 452:423–8. [PubMed: 18344981]

Fellay J, et al. A whole-genome association study of major determinants for host control of HIV-1. Science. 2007; 317:944–7. [PubMed: 17641165]

Franke A, et al. Genome-wide meta-analysis increases to 71 the number of confirmed Crohn’s disease susceptibility loci. Nat Genet. 2010; 42:1118–25. [PubMed: 21102463]

Furberg H, et al. Genome-wide meta-analyses identify multiple loci associated with smoking behavior. Nat Genet. 2010; 42:441–7. [PubMed: 20418890]

Ganesh SK, et al. Multiple loci influence erythrocyte phenotypes in the CHARGE Consortium. Nat Genet. 2009; 41:191–8. [PubMed: 19862010]

Gibbs JR, et al. Abundant quantitative trait loci exist for DNA methylation and gene expression in human brain. PLoS Genet. 2010; 6:e1000952. [PubMed: 20485568]

Greenawalt DM, et al. A survey of the genetics of stomach, liver, and adipose gene expression from a morbidly obese cohort. Genome Res. 2011; 21:1008–16. [PubMed: 21602305]

Hakonarson H, et al. A novel susceptibility locus for type 1 diabetes on Chr12q13 identified by a genome-wide association study. Diabetes. 2008; 57:1143–6. [PubMed: 18198356]

Heinzen EL, et al. Tissue-specific genetic control of splicing: implications for the study of complex traits. PLoS Biol. 2008; 6:e1. [PubMed: 1922302]

Hindorff LA, et al. Potential etiologic and functional implications of genome-wide association loci for human diseases and traits. Proc Natl Acad Sci U S A. 2009; 106:9362–7. [PubMed: 19474294]

Hoglinger GU, et al. Identification of common variants influencing risk of the tauopathy progressive supranuclear palsy. Nat Genet. 2011; 43:699–705. [PubMed: 21685912]

Kugathasan S, et al. Loci on 20q13 and 21q22 are associated with pediatric-onset inflammatory bowel disease. Nat Genet. 2008; 40:1211–5. [PubMed: 18758464]

Liu C, et al. Whole-genome association mapping of gene expression in the human prefrontal cortex. Mol Psychiatry. 2010a; 15:779–84. [PubMed: 20351726]

Liu F, et al. Digital quantification of human eye color highlights genetic association of three new loci. PLoS Genet. 2010b; 6:e1000934. [PubMed: 20463881]

McGovern DP, et al. Genome-wide association identifies multiple ulcerative colitis susceptibility loci. Nat Genet. 2010; 42:332–7. [PubMed: 20228799]

Moffatt MF, et al. Genetic variants regulating ORMDL3 expression contribute to the risk of childhood asthma. Nature. 2007; 448:470–3. [PubMed: 17611496]
Morley M, et al. Genetic analysis of genome-wide variation in human gene expression. Nature. 2004; 430:743–7. [PubMed: 15269782]

Myers AJ, et al. A survey of genetic human cortical gene expression. Nat Genet. 2007; 39:1494–9. [PubMed: 17982457]

Nalls MA, et al. Multiple loci are associated with white blood cell phenotypes. PLoS Genet. 2011a; 7:e1002113. [PubMed: 21738480]

Nalls MA, et al. Imputation of sequence variants for identification of genetic risks for Parkinson’s disease: a meta-analysis of genome-wide association studies. Lancet. 2011b; 377:641–9. [PubMed: 21292315]

Nicolae DL, et al. Trait-associated SNPs are more likely to be eQTLs: annotation to enhance discovery from GWAS. PLoS Genet. 2010; 6:e1000888. [PubMed: 20369019]

Preuss M, et al. Design of the Coronary ARtery DIsease Genome-Wide Replication And Meta-Analysis (CARDioGRAM) Study: A Genome-wide association meta-analysis involving more than 22,000 cases and 60,000 controls. Circ Cardiovasc Genet. 2010; 3:475–83. [PubMed: 20923989]

Schadt EE, et al. Mapping the genetic architecture of gene expression in human liver. PLoS Biol. 2008; 6:e107. [PubMed: 18462017]

Scott LJ, et al. A genome-wide association study of type 2 diabetes in Finns detects multiple susceptibility variants. Science. 2007; 316:1341–5. [PubMed: 17463248]

Sladek R, et al. A genome-wide association study identifies novel risk loci for type 2 diabetes. Nature. 2007; 445:881–5. [PubMed: 17293876]

Steinthorsdottir V, et al. A variant in CDKAL1 influences insulin response and risk of type 2 diabetes. Nat Genet. 2007; 39:770–5. [PubMed: 17460697]

Stranger BE, et al. Population genomics of human gene expression. Nat Genet. 2007; 39:1217–24. [PubMed: 17873874]

Tang W, et al. Genome-wide association study identifies novel loci for plasma levels of protein C: the ARIC study. Blood. 2010; 116:5032–6. [PubMed: 20802025]

Tobin JE, et al. Haplotype and gene expression implicate the MAPT region for Parkinson disease: the GenePD Study. Neurology. 2008; 71:28–34. [PubMed: 18590949]

Todd JA, et al. Robust associations of four new chromosome regions from genome-wide analyses of type 1 diabetes. Nat Genet. 2007a; 39:857–64. [PubMed: 17554260]

Todd JA, et al. Robust associations of four new chromosome regions from genome-wide analyses of type 1 diabetes. Nat Genet. 2007b; 39:857–64. [PubMed: 17554260]

Trabzuni D, et al. Quality control parameters on a large dataset of regionally dissected human control brains for whole genome expression studies. J Neurochem. 2011

Vandrovcova J, et al. Disentangling the role of the tau gene locus in sporadic tauopathies. Curr Alzheimer Res. 2010; 7:726–34. [PubMed: 20704554]

Waterworth DM, et al. Genetic variants influencing circulating lipid levels and risk of coronary artery disease. Arterioscler Thromb Vasc Biol. 2010; 30:2264–76. [PubMed: 20864672]

Webster JA, et al. Genetic control of human brain transcript expression in Alzheimer disease. Am J Hum Genet. 2009; 84:445–58. [PubMed: 19361613]

Wheeler HE, et al. Sequential use of transcriptional profiling, expression quantitative trait mapping, and gene association implicates MMP20 in human kidney aging. PLoS Genet. 2009; 5:e1000685. [PubMed: 19834535]

Wood AR, et al. Allelic heterogeneity and more detailed analyses of known loci explain additional phenotypic variation and reveal complex patterns of association. Hum Mol Genet. 2011

Wray NR, et al. Genome-wide association study of major depressive disorder: new results, meta-analysis, and lessons learned. Mol Psychiatry. 2010

Yang Q, et al. Multiple genetic loci influence serum urate levels and their relationship with gout and cardiovascular disease risk factors. Circ Cardiovasc Genet. 2010; 3:523–30. [PubMed: 20884846]

Yuan X, et al. Population-based genome-wide association studies reveal six loci influencing plasma levels of liver enzymes. Am J Hum Genet. 2008; 83:520–8. [PubMed: 18940312]
Zeggini E, et al. Meta-analysis of genome-wide association data and large-scale replication identifies additional susceptibility loci for type 2 diabetes. Nat Genet. 2008; 40:638–45. [PubMed: 18372903]
Highlights
We integrate GWAS SNPs and examine the genetic control of gene expression in blood and brain tissue.

- Many eQTLs associated with human traits are shared between blood and brain.
- A number of discrete, tissue specific eQTLs also exist in blood or brain.
- Functional studies in blood have a limited capacity to inform on regulatory variation in the brain.
- Design of eQTL mapping experiments should consider the tissue of interest for the phenotype studied.
Figure 1. Power to detect eQTLs in brain and blood
Post hoc power calculations were performed for sample sizes that we achieved after quality control in brain (A; 399 samples) or in blood (B; 501 samples). We estimated power (y axis) at a range of minor allele frequencies (x axis) for each sample series. Each colored line represents a different normalized effect size (Z) varying from 0.1 to 2.0 standard deviations of difference for each minor allele in an additive model. The steeper power curves for the blood series (B) indicate improved power over brain (A) to detect the same effect size, given the lower number of samples in the former series.
Figure 2. Comparative gene expression in blood and in brain

(A–C) Normalized gene expression values for each probe on the microarrays were converted to mean values across the population and ranked such that 1.0 is the highest expressed gene. Where genes were detected in <95% of samples in the population, we set the percentile to 0. We plotted these to compare expression in blood versus frontal cortex (A) or cerebellum (B), or to compare frontal cortex and cerebellum (C). Each probe is color coded by the difference in rank between the pairs of tissue. (D–F) Similar plot but for percentile rank of the variance in expression across the population of samples for blood versus frontal cortex (D) or cerebellum (E), or frontal cortex and cerebellum (F).
Figure 3. Similar and distinct SNP:probe associations in brain and blood
Each point shows comparisons of $-\log[10]$ of FDR corrected p-values for identical SNP and probe combinations across all 3 tissues investigated, comparing blood with frontal cortex (A) or cerebellum (B) and frontal cortex to cerebellum (C). Size of points is scaled to the combined FDR corrected p-values after $-\log[10]$ transformation. Points are colored by the associated phenotypes, where brain traits are shown in orange, blood traits in green and others in blue.
Figure 4. Blood and brain specific eQTLs in probes that are detected in all tissues
(A) Similar locus plot for ILMN_167893, which maps to ADCY3 and reveals a highly significant signal in the brain samples but no significant p values in blood, despite adequate detection of the probe in all tissues. (B) Plot of SNPs along the Chr17 region that includes the GSDML and ORMDL3 genes showing −log[10]P values for association of each SNP with expression of Illumina probe ILMN_1666206, which maps to the GSDML gene. Despite having significant detection in all three tissues, there was a strong signal for blood (red) but not in either of cerebellum (blue) or frontal cortex (green).
Figure 5. eQTLs in probes detected only in brain or blood
(A) SNPs along the region of Chr5 that contains the LRAP gene showing −log[10]P values for association of each SNP with expression of Illumina probe ILMN_1743143. (B) Plot of SNPs along the Chr17 region that includes the MAPT gene for ILMN_1710903 in frontal cortex (upper panel) or cerebellum (lower panel). For each tissue, we repeated the original eQTL analysis (green) but made the analysis conditional on a proxy SNP for the H1/H2 inversion haplotype (orange). The decrease in P values after conditioning on a proxy SNP suggests that most of the signal arises from the H1/H2 haplotype.
Table 1
Counts of SNPs and SNP:probe pairs tested and significant associations per tissue and trait for probes that were detected in all tissues

|                     | All GWAS | Blood   | Brain   | Other   |
|---------------------|----------|---------|---------|---------|
| SNPs                | 783      | 176 (22.5%) | 61 (7.8%) | 546 (69.7%) |
| SNP:probe pairs     | 2929     | 683 (23.3%) | 227 (7.6%) | 2019 (68.9%) |
| Count (% of all) of significant associations within 500kb of SNP |
| Blood               | 125      | 21 (16.8%) | 16 (12.8%) | 88 (70.4%) |
| Cerebellum          | 33       | 5 (15.15%) | 5 (15.15%) | 23 (69.7%) |
| Frontal Cortex      | 21       | 3 (14.3%) | 2 (9.5%)  | 16 (76.2%) |
Table 2
Counts of SNPs and SNP:probe pairs tested and significant associations per tissue and trait for probes that were detected in either blood or in brain.

| Tissue         | SNPs | Probes | Associations tested | All GWAS | Blood | Brain | Other |
|----------------|------|--------|---------------------|----------|-------|-------|-------|
| Blood          | 648  | 413    | 658                 | 21       | 5 (23.8%) | 2 (9.5%) | 14 (66.7%) |
| Cerebellum     | 943  | 1853   | 3924                | 107      | 21 (19.6%) | 13 (12.1%) | 73 (68.2%) |
| Frontal Cortex | 978  | 1877   | 3968                | 90       | 21 (23.3%) | 7 (7.8%) | 62 (68.9%) |