

Regulators and Psychiatric Associates of Genomic Imprinting in the Human Brain

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Abstract

We present the first genome-wide analysis of human genomic imprinting based on RNA-seq measurements of the parental bias in allele-specific expression in the dorsolateral prefrontal cortex. We find that the fraction of imprinted human genes is consistent with lower ($\approx 0.5\%$) as opposed to higher ($\approx 5\%$) estimates in mice. Our analysis reveals that age up or down-regulates allelic bias of some imprinted genes even in later life and that allelic bias depends also on gender and genetic variation. Furthermore, we show that allelic bias of some imprinted genes, like UBE3A in the Prader-Willi region, is linked to schizophrenia. These results support the hypothesized role of imprinted genes in social interactions, which dynamically change in the life course and depend on neuropsychological function.

1 Introduction

Genomic imprinting, leading to repression of either the maternal or paternal allele, has reached its highest prevalence in humans and other placental organisms [17]. In line with this, well-known physiological functions of imprinted genes include embryonic and placental development, body growth, suckling, and maternal behavior [16, 15]. Genomic imprinting requires the placement of different epigenetic marks, such as DNA methylation, at the respective alleles residing on the chromosomes originating from the mother or father [16]. Indeed, imprinted genes typically reside in clusters spanning hundreds of kilobases and allele-specific differential epigenetic marks are found near specific genes as well as in shared regulatory elements called imprinting control regions. For non-imprinted genes expression is balanced such that the two alleles are roughly equally expressed. By contrast, for imprinted genes epigenetic marks lead to substantial, degrees of imbalance in the expression level of the two alleles. We refer to

the degree of expression imbalance as *allelic bias*; at its maximum expression is completely monoallelic.

Why natural selection favors allelic bias for imprinted genes remains debated [21, 12, 11] but the most mature of all theories, kinship or conflict theory [21], provides a flexible framework for interpreting past studies on imprinted genes and formulating hypotheses and predictions regarding their detailed regulation and physiological function. The theory assumes that all imprinted genes contribute to inter-individual interactions in a highly dose-dependent fashion, and explains allelic bias with the conflicting interests of paternal and maternal genes, which arise from sexual asymmetries in those interactions [21]. A well-known asymmetry is the disproportionate role of mothers in nurturing offspring in Placentalia. Kinship theory thus explains why overexpression disorders of paternally or maternally biased genes in children abnormally promote or inhibit, respectively, their growth [16, 15].

Since different inter-individual interactions take place in various developmental stages and are mediated by various organs, kinship theory explains the non-uniform pattern of "imprintedness" and allelic bias over various ages [2] and tissue types, which is seen for several imprinted genes [16, 15]. For other genes such patterns await discovery. The theory quantitatively predicts relaxation of allelic conflict with age [20] and so raises the hypothesis that change in allelic bias is linked to aging. A study on newborn and young adult mice partially supports that hypothesis [14] but experimental evidence from humans, including older individuals, is missing.

In the framework of kinship theory the question of aging is closely tied to the roles of imprinted genes in social interactions and in the underlying psychiatric functions [20, 21], whose importance has been increasing in human evolution. Indeed, most human imprinted gene syndromes are characterized by not only growth disorder but also mental retardation and psychiatric dysfunction [16, 15]. More precisely, paternally and maternally biased genes are suggested to play antagonistic roles not only in growth but also in psychiatric functions [4], since overexpression of the former is associated with autistic while that of the latter with psychotic spectrum disorders. For example, maternally derived microduplications at 15q11-q13 may not only cause the Prader-Willi syndrome [15]—whose symptoms include obesity—but also highly penetrant for schizophrenia [10, 19], which is perhaps the most devastating psychotic spectrum disorder.

Recently the CommonMind Consortium produced and shared1 genome-wide data sets on genotype and gene expression in the dorsolateral prefrontal cortex (DLPFC) of hundreds of schizophrenic and control individuals, and identified some 650 differentially expressed genes [6]. Our present work extends that analysis with a focus on allele-specific expression across the genome, allowing us to determine allelic bias for each gene. Based on that, we find that

 $\approx 0.6\%$ of all genes are imprinted in the human DLPFC, the majority of which had been reported to be imprinted in the context of one or another tissue and/or species. We find a number of genes with allele specific expression residing near clusters of known imprinted genes. Furthermore, our data suggest that for several imprinted genes the variation of allelic bias across individuals is explained by differences in age, schizophrenia diagnosis, gender or ancestry. For example, UBE3A in 15q11-q13, appears to have allele-specific bias that varies with age and psychiatric condition.

2 Methods

2.1 Study design

We used allele-specific RNA-seq read counts from the CommonMind project to probe the allelic bias of allelic expression for each gene g that is substantially expressed in the DLPFC. We exploited the fact that single nucleotide polymorphisms (SNPs) provide a way to distinguish between the two parental alleles. For a heterozygous SNP s in a gene the reference and the alternative variant tags the maternal and paternal allele or the other way around. When allelic bias towards positively biased allele b (which may be maternal or paternal) is strong then it seems likely that the count of reads tagged by the corresponding SNP variant (which may be reference or alternative) is also relatively high.

This intuition may be formalized by introducing p, the fraction of transcripts from the parent towards which expression is biased. p is interpretable as the strength of allelic bias. Note that $1/2 \le p \le 1$. Let b identify the positively biased allele, i.e. the allele with the higher number of transcripts. Now let B denote the count of RNA-seq reads that map to the b allele at a SNP that distinguishes b from the other allele. Similarly, let T denote the total read count at that SNP. Then, assuming that B is a binomial random variable with given denominator T and expected relative frequency p, the probability that B is the higher of the two read counts, i.e. $B \ge T - B$, is

$$\sum_{x:T/2 \le x \le T} \binom{T}{x} p^x (1-p)^{T-x}.$$

The fact that this probability increases with p—as allelic bias gets stronger—supports our intuition.

These considerations motivated us to quantify allelic bias using a statistic called read count ratio S, whose definition we based on the total read count T and the higher read count <math>H, i.e. the count of reads carrying only either the reference or the alternative SNP variant,

whichever is higher. The definition is

$$S_{ig} = \frac{H_{ig}}{T_{ig}} = \frac{\sum_s H_s}{\sum_s T_s},\tag{1}$$

where *i* identifies an individual, *g* a gene, and the summation runs over all SNPs *s* for which gene *g* is heterozygous in individual *i* (Fig. 1). Note that if B_{ig} is the count or reads that map to the b_{ig} allele (defined as above) and if we make the same distributional assumption as above, namely that $B_{ig} \sim \text{Binom}(p_{ig}, T_{ig})$, then $\Pr(H_{ig} = B_{ig}|p_{ig})$, the probability of correctly assigning the reads with the higher count to the allele towards which expression is biased, tends to 1 as $p_{ig} \rightarrow 1$. We took advantage of this theoretical result in that we subjected only those genes to statistical inference, whose read count ratio was found to be high and, therefore, whose p_{ig} is expected to be high as well.

Fig. 1 illustrates the calculation of S_{ig} for the combination of two hypothetical genes, g_1, g_2 , and two individuals, i_1, i_2 . It also shows an example for the less likely event that the lower rather than the higher read count corresponds to the SNP variant tagging the higher expressed allele (see SNP s_3 in gene g_1 in individual i_2).

Using the read count ratio we analyzed two aspects of the variation of allelic bias. First, we ranked all expressed genes according to the *gene score*, a summary statistic that quantifies how right-shifted the distribution of S_{ig} is for any given gene g, and combined that information with prior evidence for imprinting. Second, given a top-scoring gene g, we sought to identify biological regulators and psychiatric consequences of allelic bias by studying the conditional distribution of S_{g} given observations $x_{1}, ..., x_{p}$ on features of study individuals that are not gene-specific. We call these features *predictors* because we used them in a regression model framework. Predictors and their various levels (if any) are listed in Table 1.

Before we carried out our read count ratio-based analyses, however, we cleaned our RNA-seq data by quality-filtering and by improving the accuracy of SNP calling with the use of DNA SNP array data and imputation. In the following subsections of Methods we describe the data, these procedures, as well as our regression models in detail.

2.2 Data

2.2.1 Brain samples, RNA-seq

Human RNA samples were collected from the dorsolateral prefrontal cortex of the Common-Mind consortium from a total of 579 individuals after quality control. Subjects included 267 control individuals, as well as 258 with schizophrenia (SCZ) and 54 with affective spectrum disorder (AFF). RNA-seq library preparation uses Ribo-Zero (which selects against ribosomal

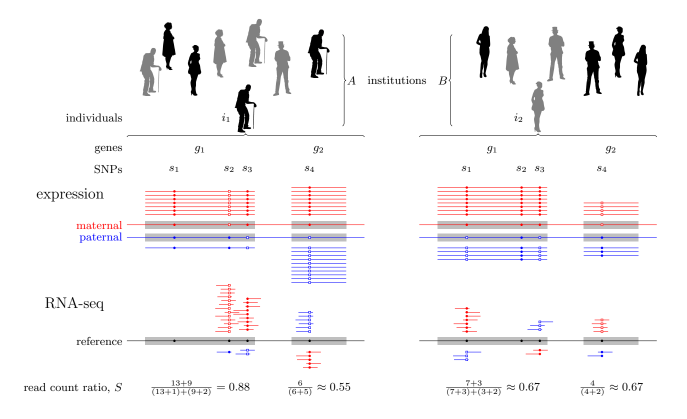


Figure 1: Quantifying allelic bias of expression in human individuals using the read count ratio statistic S_{ig} . The strength of bias towards the expression of the maternal (red) or paternal (blue) allele of a given gene g in individual i is gauged with the count of RNA-seq reads carrying the reference allele (small closed circles) and the count of reads carrying an alternative allele (open squares) at all SNPs for which the individual is heterozygous. The differences in the unobserved allelic bias between individual i_1 and i_2 arise only from biological differences such as their disease status (black and gray silhouettes), age, or gender. In addition to these, the differences in the observed read count ratio also reflect variation from technical sources like tissue preparation, or RNA sequencing. Note the sampling bias towards older individuals in institution A relative to B.

| predictor | levels |
|-------------|-----------------------------|
| Age | |
| Institution | [MSSM], Penn, Pitt |
| Gender | [Female], Male |
| PMI | |
| Dx | [AFF], Control, SCZ |
| RIN | |
| RNA_batch | [A], B, C, D, E, F, G, H, 0 |
| Ancestry.1 | |
| : | |
| Ancestry.5 | |

Table 1: Left column: predictors (explanatory variables) of read count ratio. Right column: levels of each factor-valued (i.e. categorical) predictor. Square brackets [...] surround the baseline level against which other levels are contrasted. Abbreviations: PMI: post-mortem interval; Dx: disease status; AFF: affective spectrum disorder; SCZ: schizophrenia; RIN: RNA integrity number; Ancestry.k: the k-th eigenvalue from the decomposition of genotypes indicating population structure.

RNA) to prepare the RNA, followed by Illumina paired end library generation. RNA-seq was performed on Illumina HiSeq 2000.

2.2.2 Mapping, SNP calling and filtering

We mapped 100bp, paired-end RNA-seq reads (≈ 50 million reads per sample) using Tophat to Ensembl gene transcripts of the human genome (hg19; February, 2009) with default parameters and 6 mismatches allowed per pair (200 bp total). We required both reads in a pair to be successfully mapped and we removed reads that mapped to > 1 genomic locus. Then, we removed PCR replicates using the Samtools rmdup utility; around one third of the reads mapped (which is expected, given the parameters we used and the known high repeat content of the human genome) We used Cufflinks to determine gene expression of Ensembl genes, using default parameters. Using the BCFtools utility of Samtools, we called SNPs (SNVs only, no indels). Then, we invoked a quality filter requiring a Phred score > 20 rresponding to a probability for an incorrect SNP call < 0.01).

We annotated known SNPs using dbSNP (dbSNP 138, October 2013). Considering all 579 samples, we find 936,193 SNPs in total, 563,427 (60%) of which are novel. Further filtering of

this SNP list removed the novel SNPs and removed SNPs that either did not match the alleles reported in dbSNP or had more than 2 alleles in dbSNP. We also removed SNPs without at least 10 mapped reads in at least one sample. Read depth was measured using the Samtools Pileup utility. After these filters were applied, 364,509 SNPs remained in 22,254 gene These filters enabled use of data with low coverage. For the 579 samples there were 203 million reads overlapping one of the 364,509 SNPs defined above. Of those 158 million (78%) had genotype data available from either SNP array or imputation.

2.2.3 Genotyping and calibration of imputed SNPs

DNA samples were genotyped using the Illumina Infinium SNP array. We used PLINK with default parameters to impute genotypes for SNPs not present on the Infinium SNP array using 1000 genomes data. We calibrated the imputation parameters to find a reasonable balance between the number of genes assessable for allelic bias and the number false positive calls since the latter can arise if a SNP is incorrectly called heterozygous.

We first examined how many SNPs were heterozygous in DNA calls and had a discordant RNA call (i.e. homozygous SNP call from RNA-seq) using different imputation parameters. Known imprinted genes were excluded. We examined RNA-seq reads overlapping array-called heterozygous SNPs which we assigned a heterozygosity score $L_{\rm het}$ of 1, separately from RNA seq data overlapping imputed heterozygous SNPs, where the $L_{\rm het}$ score could range from 0 to 1. After testing different thresholds we selected an $L_{\rm het}$ cutoff of 0.95 (i.e. imputation confidence level of 95%), and a minimal coverage of 7 reads per SNP. With these parameters, the discordance rate (monoallelic RNA genotype in the context of a heterozygous DNA genotype) was 0.71% for array-called SNPs and 3.2% for imputed SNPs.

The higher rate of discordance for the imputed SNPs is due to imputation error. These were taken into account in two ways. First, we considered all imputed SNPs for a gene g and individual i jointly. Second, we excluded any individual, for which one or more SNPs supported biallelic expression.

2.2.4 Quality filtering

Two kind of data filters were applied sequentially: (1) a read count-based and (2) an individual-based. The read count-based filter removes any such pair (i, g) of individual i and genes g for which the total read count $T_{ig} < t_{rc}$, where the read count threshold t_{rc} was set to 15. The individual-based filter removes any genes g (across all individuals) if read count data involving g are available for less than t_{ind} number of individuals, set to 25. These final filtering procedures decreased the number of genes in the data from 15584 to n = 5307.

2.3 Statistical analysis

2.3.1 Test for nearly unbiased expression

This test was defined by the criterion

$$S_{ig} \le 0.6 \text{ and } UCL_{ig} \le 0.7,$$
 (2)

where the 95% upper confidence limit UCL_{ig} for the expected read count ratio p_{ig} was calculated based on the assumption that the higher read count $H_{ig} = S_{ig}T_{ig} \sim \text{Binom}(p_{ig}, T_{ig})$, on the fact that binomial random variables are asymptotically (as $T_{ig} \to \infty$) normal with $var(H_{ig}) = T_{ig}p_{ig}(1 - p_{ig})$, and on the equalities $var(S_{ig}) = var(H_{ig}/T_{ig}) = var(H_{ig})/T_{ig}^2$. Therefore

$$UCL_{ig} = S_{ig} + z_{0.975} \sqrt{\frac{S_{ig}(1 - S_{ig})}{T_{ig}}},$$
(3)

where z_p is the p quantile of the standard normal distribution.

2.3.2 Regression models

Let m denote the number of individuals/samples and \mathcal{G} the set of n = 5307 genes that passed quality filtering. Regression analysis involved a subset $\mathcal{G}_1 \subset \mathcal{G}$ of $n_1 = 30$ genes called as imprinted.

The basic model, unlm.S, is

$$\mathbf{S} = \mathbf{X}\boldsymbol{\beta} + \boldsymbol{\varepsilon},\tag{4}$$

$$\varepsilon_{ig} \stackrel{\text{iid}}{\sim} \text{Norm}(0, \sigma_q^2)$$
 (5)

where the response **S** is an $m \times n_1$ read count matrix with entries S_{ig} , X is an $m \times p$ design matrix, $\boldsymbol{\beta}$ is a $p \times n_1$ matrix of regression coefficients (Table 1), the random error $\boldsymbol{\varepsilon}$ has the same dimension as **S**, and gene $g \in \mathcal{G}_1$. Eq. 4 may be given as

$$S_g = \mathbf{X}\beta_g + \varepsilon_g,\tag{6}$$

where the vectors S_g , β_g , ε_g are single columns taken from their respective matrix counterparts. The unlm.S model was extended in several ways, yielding

- 1. six normal linear models (Table 2)
- 2. two logistic models logi. S and logi2. S.

| | weights w_{ig} | |
|-------------------|------------------|----------|
| | 1 | T_{ig} |
| no transformation | unlm.S | wnlm.S |
| quasi-log transf. | unlm.Q | wnlm.Q |
| rank transf. | unlm.R | wnlm.R |

Table 2: Specification and notation of normal linear models based on the weight w_{ig} on each observation and the transformation τ applied to the set $\{S_{ig}|g\}$ of read count ratios for a given gene g.

The general form of the normal linar models (cf. 6) is

$$\mathbf{W}_g^{1/2}\tau(S_g) = \mathbf{W}_g^{1/2}\mathbf{X}\beta_g + \varepsilon_g. \tag{7}$$

The extension here consists of \mathbf{W}_g , an $m \times m$ diagonal matrix of weights w_{ig} on the *i*-th diagonal position, and τ , a transformation on read counts. Besides the trivial identity transformation (i.e. no transformation) two kinds of transformation were used: the rank transformation and a quasi-log transformation τ_Q defined as

$$\tau_Q(S_{ig}; T_{ig}) \equiv Q_{ig} = -\log\left(1 - S_{ig}\frac{T_{ig}}{T_{ig} + c}\right),\tag{8}$$

where log means natural logarithm (base e). c is a pseudo read count set to 1 in order to avoid zero in the parenthesis since the log function is undefined at 0.

The logistic models, logi. S or logi2. S, share the general form

$$S_a = \mu_a + c \,\varepsilon_a \tag{9}$$

$$\mu_g = h(\mathbf{X}\beta_g) \tag{10}$$

$$\varepsilon_{ig} + \mu_g \stackrel{\text{iid}}{\sim} \text{Binom}(\mu_g, T_{ig}).$$
 (11)

The link function h is $h(u) = e^u/(1 + e^u)$ for logi.S and $h(u) = e^u/(2 + 2e^u)] + 1/2$ for logi2.S, and the scaling constant c is 1 and 1/2, respectively. Thus, the response S_g under logi2.S is scaled and shifted relative to that under logi.S such that (with probability one) $1/2 \le S_{ig} \le 1$ under the former and $0 \le S_{ig} \le 1$ under the latter.

Each of the eight models (including normal linear and logisite models) has $p \times n_1$ regression parameters corresponding to the dimension of $\boldsymbol{\beta}$. This allows different behavior for different genes since $\beta_1 \neq ... \neq \beta_{n_1}$ in general. Therefore, the estimated regression coefficients are reported as $\hat{\beta}_g = (\hat{\beta}_{1g}, ..., \hat{\beta}_{jg}, ..., \hat{\beta}_{pg})$ for each gene g, often replacing index j with the name of the parameter such as Age or InstitutionPitt (Table 1).

A second set of eight models was also fitted, for which β was constrained such that $\beta_1 = ... = \beta_{n_1}$. This was achieved by aggregating over genes $g \in \mathcal{G}_1$ the higher read count $H'_i = \sum_g H_{ig}$, the total read count $T'_i = \sum_g T_{ig}$, redefining the read count ratio as $S'_i = H'_i/T'_i$, and replacing S_g by $S' = (S'_1, ..., S'_m)$ in Eq. 6, 7, 9, and T_{ig} by T'_i in Table 2 and Eq. 11. Note that such aggregation simplifies the matrix variables in Eq. 4 to the corresponding vector variables in Eq. 6. Because S'_i is a weighted average of $\{S_{ig}\}_i$, results under these models are reported as $\hat{\beta}_{WA} = (\hat{\beta}_{1WA}, ..., \hat{\beta}_{jWA}, ..., \hat{\beta}_{pWA})$.

These 2×8 models are all multiple regression ones with p < 1 parameters. Two corresponding sets of models with a single Age parameter (p = 1) were also fitted but the results were only used for visual illustration of model fits in Fig. 5 but not for quantitative inference.

3 Results

3.1 Genome- and population-wide variation of allelic bias

A total of 5307 genes passed our filters designed to remove genes with scarce RNA-seq data reflecting low expression and/or low coverage of RNA-seq. Examining these genes, we performed exploratory statistical analysis based on the read count ratio statistic S_{ig} , whose results (below) we interpreted in terms of the variation of allelic bias both across genes g and across individuals i.

Fig. 2 presents the conditional empirical distribution of S_{g} given each gene g. Each of the three plots of the upper half show in a distinct representation the same empirical distributions based on data for three genes. The main panels of the lower half present, in the most compact representation, the distributions based on all data (5307 genes). Two of the three genes in the upper half, PEG10 and ZNF331, are known imprinted genes in the sense that they had previously been found imprinted in the context of some developmental stage, species, and tissue type other than the adult human DLPFC. The third, AFAP1, is referred to as a candidate (imprinted) gene as it has not been reported to be imprinted in any context. For all three genes S_{g} varies considerably within its theoretical range $\left[\frac{1}{2}, 1\right]$. This suggests variation of allelic bias across study individuals, although some component of the variation of S_{g} must originate from technical sources. Later subsections present our modeling and detailed analysis of the across-individual variation on genes called imprinted in the context of adult human DLPFC. The rest of the present subsection reports the calling of imprinted genes.

We defined gene score as the location statistic $1 - \text{ECDF}_g(0.9)$, the fraction of individuals i for which $S_{ig} > 0.9$. This score is shown in side plots of the lower half of Fig. 2 as gray filled circles or, for the three genes mentioned above, as larger green circles (the latter are also present

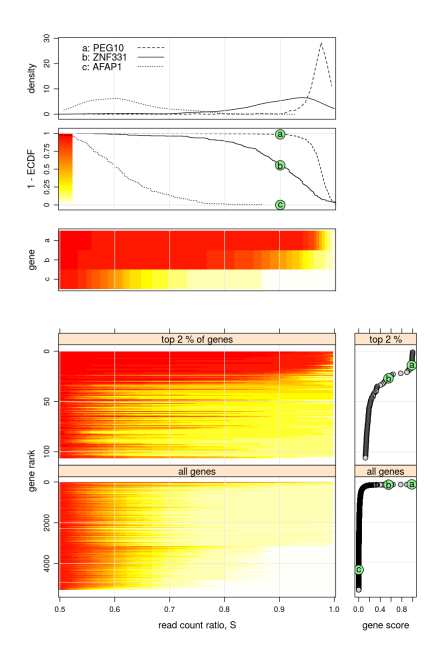


Figure 2: Using the read count ratio statistic S to report on variation of allelic bias across individuals and genes. $Upper\ half$, from top to bottom: (1) kernel density estimate; (2) the graph of the survival function 1 - ECDF, where ECDF means empirical cumulative distribution function; note color scale for heat map and green filled circles marking genes' score; (3) the heat map representation of the survival function. $Lower\ half$, main panels: heat map of the survival function for all 5307 analyzed genes ranked according to their score; right side panels: gene score.

in the second from top graph). Based on the score genes were ranked; the heat map of empirical distribution of S_{g} of ranked genes suggests that the top 50 genes, which constitute $\approx 1\%$ of all genes in our analysis, are qualitatively different from the bottom $\approx 99\%$ suggesting that most of them are imprinted. Consistent with this, the top-scoring genes tended to cluster around genomic locations that had been previously described as imprinted gene clusters (Fig. S1).

The set of top scoring 50 genes is highly enriched in known imprinted genes, marked by blue in Fig. 3 and in nearby candidate genes (green) defined as being within 1Mb of a known imprinted gene. Within the top 50, we find 29 such genes; 21 known imprinted genes and eight nearby candidates. In subsequent analysis (below) we also consider UBE3A, which falls outside of the top 50 as demonstrating allelic bias consistent with imprinting in the context of human adult DLFPC (Fig. S2).

The remaining 21 genes in the top 50 are separated by > 1 Mb from some known imprinted gene (termed distant candidates, red in Fig. 3). Upon further examination these distant candidate genes are overwhelmingly likely not imprinted. The primary reason for this conclusion is that we performed a test to see if there is reference allele bias for all candidate genes. For any gene (known imprinted, or candidate) the expectation is that when some allelic bias is detected, that should equally favor the reference or non-reference allele since for a given individual who is heterozygous at a given SNP in the genome it is reasonable to assume that the chances are equal that the mother or that the father has the reference allele. Most known imprinted genes and the nearby candidates display a reference/non-reference distribution consistent with a binomial distribution with a probability of 0.5 for both the reference and non-reference alleles. However, and in sharp distinction, most distant candidates have distributions of reference/non-reference that are not consistent with equal probabilities (see genes marked with "X" in Fig. 3). Indeed, for most of them the distribution is shifted towards the reference allele strongly suggesting that mistaken genotyping, imputation or a mapping issue led to the presence of these red genes in the list of the top 50 genes. One could argue that we should have left these genes out of Fig. 3, but we thought it was important to show them and to indicate the reasons they are set aside. Note also, that we also tested the hypothesis for each gene g and individual i that allelic expression is (nearly) unbiased (Eq. 2). The fraction of individuals for which the test was not rejected tends to be much higher for the "red" genes in the top 50 (black bars in Fig. 3).

While the shifted distribution of reference/non-reference alleles leads us to discount the possibility of imprinting, random monoallelic expression is still a distinct possibility for these candidates as our studies of random monoallelic expression in mice suggested that a substantial fraction (40 - 80%) of random monoallelically-expressed genes had a very strong bias towards monoallelic expression of one of the two alleles [23]. Moreover, it is worth noting that three

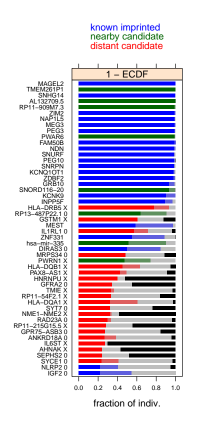


Figure 3: The top 50 genes ranked by the gene score. The score of gene g is $1 - \text{ECDF}_g(0.9)$, the fraction of individuals i for which $S_{ig} > 0.9$ and is indicated by the length of dark blue, dark green or dark red bars. Note that the same ranking and score is shown in the bottom half of Fig. 2. The right border of the light blue, light green and light red bars is at $1 - \text{ECDF}_g(0.8)$. The length of the black bars indicates the fraction of individuals passing the test of nearly unbiased expression (Eq. 2). "X" characters next to gene names indicate reference allele bias, while "0" indicate that reference allele bias could not be determined due to small amount of data.

of these candidates are from the major histocompatibility locus (HLA), which is notable for extensive polymorphism and difficulties with allelic identification. For these three genes we also analyzed them more thoroughly with HLA-specific methods for determining haplotype based on RNA-seq [1] and genotype data [22]. The high observed read count ratios for HLA genes appear to be driven by eQTL-like effects, not by random monoallelic expression nor by imprinting (manuscript in preparation). Examining all the assessable known imprinted genes, we find than $\frac{1}{3}$ rd of them have a low gene score. This suggests that these genes do not display imprinted expression in the human adult DLPFC, consistent with many reports in the literature indicating that known imprinted genes are often imprinted in some but not all tissues.

3.2 Selection among predictive models of allelic bias

We studied further the sources and psychological consequences of the variation of allelic bias in 30 genes that the above results suggested to be imprinted in the human adult DLPFC. These 30 genes consist of the 29 known imprinted or nearby candidates in the top 50 (Fig. 3) as well as UBE3A (Fig. S2). For these genes we characterized in detail the dependence of the read count ratio on biological and technical explanatory variables referred to as predictors (Table 1).

Fig. 4 shows the pattern of dependence of the read count ratio S_{g} for a given gene g on age and gender. From this visual inspection it seems that for several genes age is informative to the distribution of S_{g} in terms of both the location (e.g. the mean of S_{g}) and scale (e.g. variance); such apparent dependence on gender is not clear.

This qualitative result, however, is greatly complicated by the association among predictors: taking only pairwise associations the situation is already complex given the observed strong association between age and gender with each other and with many other predictors (Fig. S3) but higher order associations might also exist in the data. The correct interpretation of plots like Fig. 4 also depends on the amount of data, i.e. the total read count T_{ig} , based on which the read count ratio S_{ig} was calculated. Fig. S4 shows how T_{ig} varies both within a gene and across genes.

These results motivated us to model the dependence of read count ratio for a given gene on all predictors jointly in a multiple regression framework. More specifically, we used generalized linear models (GLMs). With GLMs we sought to find a reasonable compromise between simplicity and generality. While that simplicity facilitates parameter estimation and tests for independence, generality allows fitting several GLM families, among which the best family or families can then be selected based on the fit. Thus the purpose of fitting was twofold: (1)

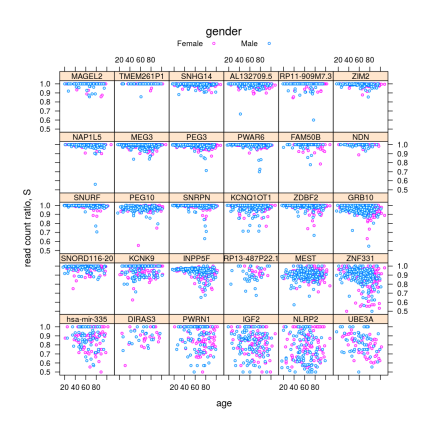


Figure 4: Variation of the read count ratio S_{ig} across hundreds of individuals i (dots) and 30 genes g that have been called imprinted in the DLPFC brain area in this study.

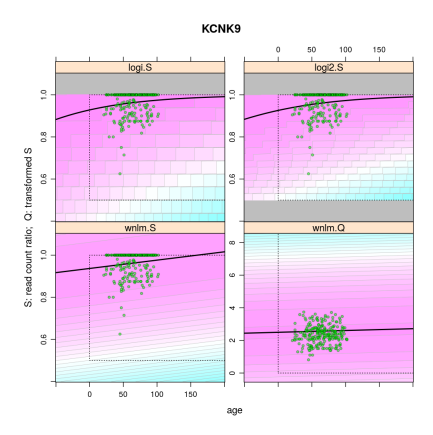


Figure 5: Fitting four families of generalized linear models to the same data set. In each panel the horizontal axis is age and the vertical axis is the possibly transformed read count ratio S. For all panels the green scatter represents the same data set (KCNK9 gene, cf. Fig4) except that in the bottom right panel the observed read count ratio S is transformed to Q according to Eq. 8. Given each one of four model families the predicted read count ratio is represented by thick black curves. The probability mass or density of the sampling distribution is indicated by a cyan-to-magenta gradient and various prediction limits by gray lines. Note that for demonstration the depicted fitted models are all simple in the sense that they contain age as a sole predictor. Quantitative inference, however, was based on the corresponding multiple regression models including all predictors (Table 1).

model selection, as well as (2) inference of dependencies between the read count ratio and predictors given the selected model family or families.

Out of the eight GLM families two were logistic (logi.S, logi2.S in Fig. 5 top). These have several desirable theoretical properties for count-based data: they give zero probability for values $S_{\cdot g} > 1$, they account for the observed dependence of the variance of $S_{\cdot g}$ on its mean, and take into account the observed total read counts. These properties follow from the fact that logistic models are natural extensions of binomial models conditioned on the observed total read count T_{ig} . The remaining six fitted GLM families contain various normal linear models (see Fig. 5 bottom for two of these). Although these are in general less suitable for count-type data, they are robust and easily interpretable. The families of normal linear models of this study are characterized by two features (Table 2): (i.) whether or not they are weighted by T_{ig} and (ii.) the transformation, if any, that had been applied to S_{ig} before the fit. Transformation proved critical (see below) but weighting had little impact either on model fit or parameter estimates (not shown) so we removed unweighted models (unlm.Q, unlm.R, unlm.S) reasoning that the mentioned small differences under weighted models reflect their improved power due to their ability to utilize information in the observed total read count.

We addressed model fit by using both Akaike Information Criterion (AIC) and diagnostic graphs based on standardized deviance residuals. However, only the latter approach turned out conclusive because the estimate for AIC is more biased for certain GLM families than for others, which rendered comparison based on the information criterion unreliable for this problem. In particular, the homoscedasticity (constant error variance) assumed by normal linear models is severely violated for wnlm.S because of the strong scale-location dependence of the read count ratio S (Fig. 4). As a consequence of that dependence, not only the location but also the scale of untransformed S depended on age, and thereby prevented good fit of the normal linear model wnlm.S (bottom left panel of Fig. 5). This problem was overcome by a quasi-log transformation Q of S (Eq. 8) that selectively abolished the dependence of the scale (bottom right panel of Fig. 5). The wnlm.Q model fits the data well for all 30 genes, as evidenced by three distinct diagnostics based on residuals (Fig. S5, S7, S9). Rank transformation R and the fitting of wnlm.R lead to a much lesser improvement than Q and wnml.Q (not shown).

The same model checking diagnostics for the logistic models suggested good fit under logi.S for eight of the 30 genes (Fig. S6, S8, S10). These genes are highlighted with bold font in the top of Fig. 6. For subsequent analysis we selected the wnlm.Q model for all 30 genes, and additionally considered the logi.S model for the eight genes with good fit.

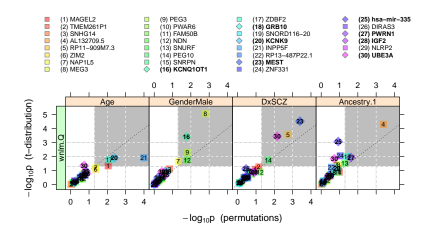


Figure 6: Significance of association between biological predictors and imprinted genes in the DLPFC. Association was tested based on the p-value for the null hypothesis $\beta_{jg} = 0$, where the regression coefficient β_{jg} mediates the dependence of the read count ratio for gene g on predictor/"treatment" j (Age,...) in the wnlm.Q model (Fig. 5 bottom right). Note that the p-values estimated parametrically (t-distribution) agree well with those estimated non-parametrically (random permutations). The gray area shows the decision rule for rejecting the null hypothesis and thus contains significant associations.

3.3 Biological predictors and interpretation

We selected four biological terms in our linear predictor (Eq. 7, 10). Of these terms Age and Ancestry.1 represent the corresponding continuous predictors whereas GenderMale and DxSCZ are treatment levels of the corresponding categorical predictors (Gender and Dx, respectively) contrasted with the corresponding control levels (GenderFemale and DxControl, see Table 1). We left out DxAFF because of the small number of AFF individuals appeared to substantially decrease the power of the hypothesis test described below.

For each term j, and each gene g called imprinted in the present context, we tested the null hypothesis $\mathcal{H}_{jg}^0: \beta_{jg} = 0$ meaning that the read count ratio is independent of that term (Fig 6). The probabilistic interpretation of the rejection of \mathcal{H}_{jg}^0 is simply that allelic bias depends on the predictor. The mechanistic interpretation is more complicated because such probabilistic dependence may manifest from either one of the following causal relations: (1) the predictor regulates the allelic bias of the gene; (2) allelic bias regulates the predictor; (3) they both regulate or are both regulated by a third, latent, variable.

Given a model family (wnlm.Q or possibly logi.S) each hypothesis was tested both parametrically and non-parametrically (see axes labeled with "t-distribution" and "permutations",

| Gene | Gene type | Chr | Coefficient | Known phenotype |
|--------------|----------------|----------------------|---------------------|---|
| ZDBF2 | protein coding | 2 | Age, Ancestry.1 | |
| NAP1L5 | protein coding | 4 | ${\bf Gender Male}$ | |
| PEG10 | protein coding | 7 | DxSCZ | |
| MEST | protein coding | 7 | DxSCZ | Silver-Russell syndrome |
| KCNK9 | protein coding | 8 | Age | Birk-Barel mental retardation dysmorphism syndrome |
| INPP5F | protein coding | 10 | Age | cell motility; endocytic recycling |
| KCNQ1OT1 | antisense | 11 | ${\bf Gender Male}$ | Beckwith-Wiedemann syn.; Isol. hemihyperplasia |
| MEG3 | lincRNA | 14 | ${\bf Gender Male}$ | Mat/pat 14q32.2 hypermeth/microdel syndrome |
| RP11-909M7.3 | lincRNA | 14 | DxSCZ | |
| AL132709.5 | miRNA | 14 | Ancestry.1 | |
| MAGEL2 | protein coding | 15 | Age | Prader-Willi syn.; Schaaf-Yang syn.; Arthrogryposis |
| NDN | protein coding | 15 | ${\bf Gender Male}$ | Prader-Willi syndrome |
| PWRN1 | lincRNA | 15 | Ancestry.1 | Prader-Willi syndrome |
| UBE3A | protein coding | 15 | DxSCZ | Prader-Willi syn.; Angelman syn.; circadian rhythm |
| PEG3 | protein coding | 19 | ${\bf Gender Male}$ | |

Table 3: Properties of genes with significance of association to one or more biological predictors.

respectively, on Fig. 6, S12 and S11). The p-value estimates show good agreement across estimation methods (parametric vs. non-parametric, Fig. S12). This agreement is expected from the regularity of the likelihood function (Fig. S14), since that allows the parametric method—which is based on asymptotic properties of maximum likelihood estimation in the limit of infinite sample size—to work well already at the current sample size. The agreement across models was weaker but still acceptable (Fig. S12); the test under logi.S appeared more powerful and/or more biased than that under wnlm.Q for most cases. Using a heuristic decision rule (gray area in Fig. 6) we rejected the marginal hypothesis \mathcal{H}_{jg}^0 if the conditional hypotheses " \mathcal{H}_{jg}^0 parametric method" and " \mathcal{H}_{jg}^0 non-parametric method" are both rejected at size $\alpha = 0.05$ under the wnlm.Q model.

For each of the four terms (Age,...) the hypothesis of independence is rejected for some genes (Fig. 6). This appears to suggest that aging, gender and ancestry-related genetic variation regulates allelic bias of those genes whereas the risk of schizophrenia is regulated by it; but, as discussed above, indirect causation involving a latent variable is also possible.

Table 3 presents some known properties of all genes for which read count ratio appears to depend on at least one of the four biological terms. For some of these genes the dependence is consistent with the prior information on the gene's physiological or pathological role. For instance, association of UBE3A to schizophrenia is consistent with its suggested role as a risk factor for this and other psychiatric conditions based on variation of its gene dose and copy number variation of its broader locus 15q11q13 (PraderWilli/Angelman region) [19, 13]. For

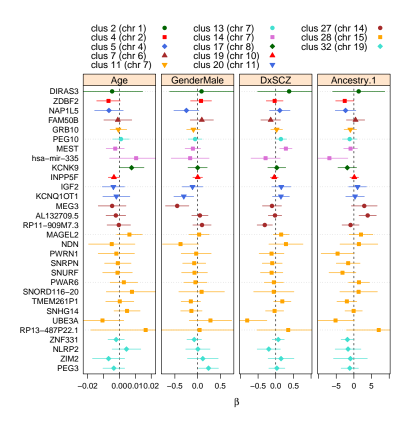


Figure 7: Dependence of allelic bias for each imprinted gene g on each biological predictor/term j quantified by the regression coefficient $\beta_{j} = 0$. The null hypothesis of no dependence is marked by the dashed vertical line at $\beta_{jg} = 0$. Each estimate $\hat{\beta}_{jg}$ is represented by a filled symbol and each 99% confidence interval by a horizontal bar.

another example, PEG3, see the Discussion.

Fig. 7 and S13 extend the above results in two ways. First, they present the maximum likelihood estimates $\hat{\beta}_{jg}$ as colored symbols, which are to be compared to the dashed lines that mark the theoretical value $\beta_{jg} = 0$ under the null hypothesis. For each regression term j the $\hat{\beta}_{jg}$ under wnlm.Q agreed reasonably with that under logi.S (Fig. S15). The estimates $\hat{\beta}_{jg}$ inform on the direction and size of the effect of each term j on read count ratio of each significantly affected gene g. Along with these quantities the 99% confidence intervals are also presented (horizontal bars in Fig. 7 and S13; see also Fig. S14 for the illustration of the link between confidence intervals and the geometry of the log-likelihood surface). For any given term we observe statistically significantly affected genes with both negative (e.g. INPP5F) and positive (e.g. KNCK9) direction. For instance, the read count ratio of INPP5F depended negatively on age, while that of KCNK9 depends on age positively, suggesting that aging may both down-

and up-regulate allelic bias in general. Interestingly, we find that both genes for which the dependence on DxSCZ is negative (UBE3A and RP11-909M7.3/MEG8) had been previously established as maternally expressed, whereas both genes with positive dependence (PEG10, MEST) as paternally expressed.

Of note is that the preceding analysis of overall expression based on the CommonMind data [6] found only PEG10 and IGF2 to be differentially expressed in schizophrenia out of the 30 genes our present analysis called imprinted in the DLPFC (Fig. S23). PEG10, but not IGF2, is among the 4 genes whose allelic bias, rather than overall expression, is significantly associated with schizophrenia.

The second extension in Fig. 7 and S13 is the arrangement of genes according to their chromosomal location and containment in various imprinted gene clusters. We find clusters (e.g. clus 14 on chr 7, clus 27 on chr 14, and clus 28 on chr 15) within which some gene exhibit significant dependence on a given predictor while all other genes did not. For a presumably causal predictor such as age this result might mean that if the significant dependence indeed reflects true association between that gene and the predictor, then all other genes in the cluster are either independent or dependent only to a degree that could not be detected as significant change. However, we do not observe two significant dependencies of opposing direction in the same cluster, which was expected given the central role of of a single imprinting control region in the establishment and maintenance of imprinting and allelic bias for all genes in a cluster.

3.4 Methodological limitations

The previous section focussed primarily on the question of dependence between a gene's read count ratio and a predictor. Therein $\hat{\beta}_{jg_1}, \hat{\beta}_{jg_2}, ...$ were compared (Fig. 7) to provide relative effect sizes across genes $g_1, g_2, ...$ for a given predictor j. But how do predictors compare to each other for any given gene? In particular, of the total variation in read count ratio what fraction is explained by technical, and what fraction by biological predictors? Can we estimate the variation in allelic bias by removing the technical variation from all explained variation of read count ratio? Unfortunately these questions proved unresolvable within the present experimental design and fitted models. We performed ANOVA with the aim of assigning a component of variation to each predictor but this failed because the reduction of residual sum of squares on the addition of a predictor to the model strongly depended on the sequence in which predictors were added (see Fig. S22 for a forward and reverse sequence). Such failure of ANOVA follows from the non-orthogonality of terms (predictor variables and their levels) in the linear predictor due to the dependencies among predictors (Fig. S3).

We observed non-orthogonality not only among terms in the linear predictor but also

among the regression coefficients (Fig. S19). This means that the log-likelihood ℓ of a given coefficient, such as β_{Age} depended on other coefficients. The dependence of $\ell(\beta_{Age})$ on β_{RIN} , the coefficient for RNA integrity number, appeared particularly strong pointing to the sensitivity of our analysis to RNA quality.

It is reasonable to assume for several predictors that their effects on the read count ratio are not, or only weakly, specific to any gene. Such predictors are RIN and the ones identifying RNA batches. Accounting for both gene specific and unspecific effects at the same time cannot be achieved in the present GLM framework (Fig. S16 left); therefore all fitted GLM families mentioned so far assumed gene specific effect for all predictors. Both under wnlm.Q and under logi.S RIN and the RNA_batchB,...RNA_batch0 predictors were found to differentially affect the read count ratio for different genes (Fig. S17, S18). This result is hard to explain mechanistically. Rather, it seems to highlight the mentioned limitation of the GLM framework. One consequence of this limitation is that, due to the discussed non-orthogonality of parameters, some technical effects may manifest as biological ones and vice versa.

Another consequence of the lack of information sharing is relatively low power. More complex models of this characteristic would have even lower power but complexity may be needed to account for possible interactions among predictors in general. To address interactions for the present data we studied the contextual dependence of read count ratio on age (Fig. S20, S21). In the context of different institutions or genders the estimates for β_{Age} were somewhat different from the marginal estimates $\hat{\beta}_{Age}$ seen in Fig. 7 and S13. This suggests that interactions are indeed present between age and some other predictors but the identity of these predictors is uncertain due to the unknown structure of dependencies among them.

We also fitted model families (Fig. S16 middle) that assume that the effects of all predictors are *not* gene specific and so pool even heterogeneous information together across genes. But this approach did not find significant dependence of read count ratio on any of the biological predictors (Fig. S17), which is consistent with the our earlier result (Fig. 7) that each of biological predictor affected relatively few genes significantly and these effects were often of opposite direction.

4 Discussion



We present the first genome-wide characterization of allelic bias of expression in humans and, at the same time, the first genome-scale study of the potential for imprinted genes' impact on a psychiatric disorder. Important to mention first, but not surprising, is our finding that $\approx \frac{2}{3}$ rd of known imprinted genes display imprinting in the adult human DLPFC. The fact that the remaining $\approx \frac{1}{3}$ rd of known imprinted genes do not display imprinting is consistent with

many other studies showing that imprinted genes often display imprinting in some but not all tissues. We also find allelic bias consistent with imprinting for eight novel candidate genes, all of which are nearby known imprinted genes. Strong allelic bias for $\approx 0.6\%$ of all assessable genes suggests that they are imprinted, which agrees closely with the most recent mouse studies [5, 14] but contradicts a controversial earlier estimate of $\approx 5\%$ also from mouse [7].

We also examined the dependence of allelic bias on variables such as age, diagnosis, gender and ancestry. We comment on a number of intriguing findings with respect to the impact of these variables in the next few paragraphs. One must bear in mind, however, that the biological interpretation of the findings depends on the correctness of our statistical approach. Although we did perform model selection based on model fit, we found that even the best fitting generalized linear models had several limitations (Section 3.4). Using an approach that conservatively combined parametric and non-parametric hypothesis testing, we find several imprinted genes for which allelic bias is dependent on aging and/or schizophrenia. These findings are in line with the general prediction from kinship theory that the roles of imprinted genes may dynamically change even at later stages of life in order to mediate certain social interactions that are dependent on psychology and cognition [20, 21].

If we accept that the main effect of allelic bias is exerted on overall expression level, our finding that the allelic bias for UBE3A is negatively correlated with schizophrenia implies that it is its overexpression that increases risk for the disorder. This agrees with the previously observed effect of copy number variation of 15q11-q13, the Prader-Willi region [13]. Moreover, we find that another maternally biased gene, RP11-909M7.3, is also negatively correlated with schizophrenia, while two paternally biased genes, MEST and PEG10, are both positively correlated. This is exactly the pattern that is predicted by kinship theory, more precisely by its special case, the imprinted brain theory [4].

The interpretation of our results on ancestry and gender is less straight-forward than that to age and schizophrenia. What appears clear, though, is that dependence on ancestry indicates various effects of genetic polymorphisms. Methylation QTLs represent one class of such effects and they might be particularly important because altering methylation at imprinting control regions can potentially regulate multiple imprinted genes within the same cluster. What's more, methylation QTLs were found to be enriched in schizophrenia risk loci [9] even though imprinting can make it more difficult to find methylation QTLs. Therefore, it will be worth investigating the methylation link between polymorphisms and allelic bias in combined genomic experiments. Dependence on gender may indicate sex-dependent physiological and behavioral functions. Our finding on PEG3 appears to be consistent with its observed involvement in sexual dimorphism in the brain and in sexual and maternal behavior [3]. The latter observations motivated the hypothesis that imprinting follows from mother-offspring

In conclusion, our analyses of allele-specific expression in the human DLPFC reveal some new candidate imprinting genes, while supporting the emerging idea that the brain is similar to other tissues in terms of the overall genome-wide extent of imprinting. We also find variation in allelic bias across individuals for all analyzed imprinted genes, and our statistical inference suggests that for several genes that variation is explained in part by age, psychiatric diagnosis, gender and ancestry. Further studies of other samples and brain regions will allow continued exploration of the impact of allele-specific gene regulation on normal and dysfunctional traits of human psychology and cognition.

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5 Supplementary Material

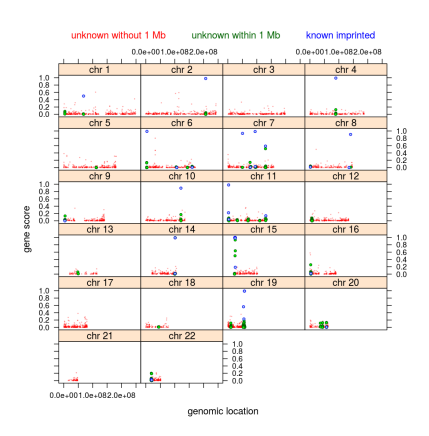


Figure S1: Clustering of top-scoring genes in the context of human DLPFC around genomic locations that had been previously described as imprinted gene clusters in other contexts.

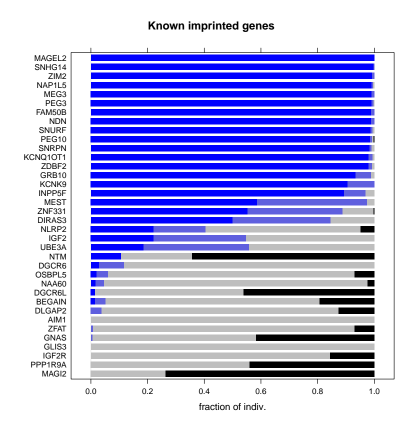


Figure S2: Known imprinted genes ranked by the gene score (dark blue bars). "Known imprinted" refers to prior studies on imprinting in the context of any tissue and developmental stage. The length of the black bars indicates the fraction of individuals passing the test of nearly unbiased expression.

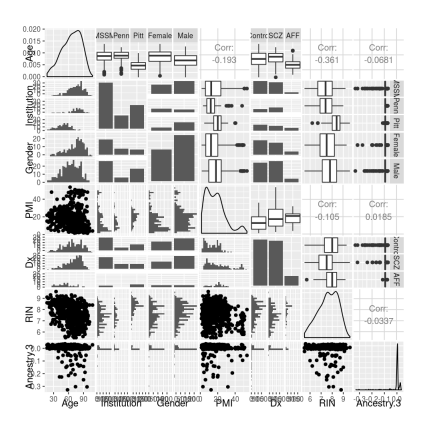


Figure S3: Pairwise dependencies among predictors.

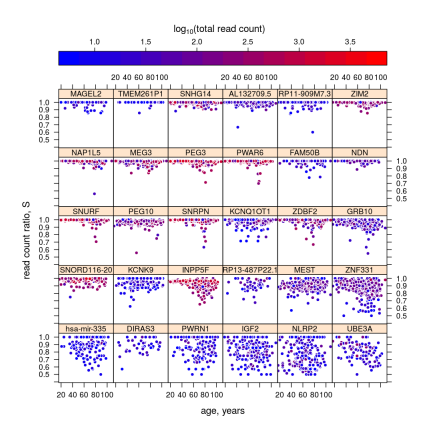


Figure S4: Variation of total RNA-seq read count across genes and individuals.

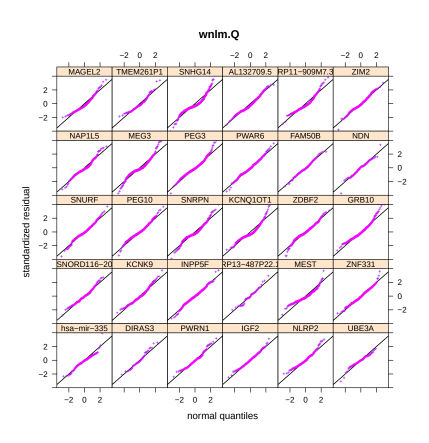


Figure S5: Checking the fit of wnlm.Q model: analysis of the normality of residuals.

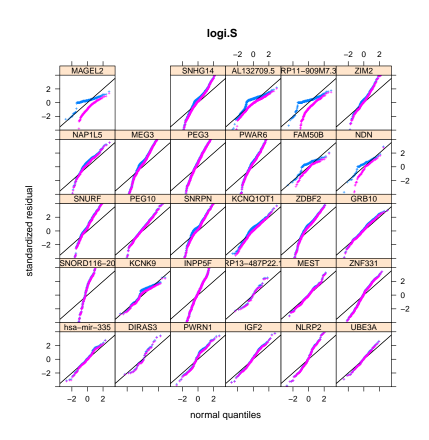


Figure S6: Checking the fit of logi.S model: analysis of the normality of standardized deviance residuals.

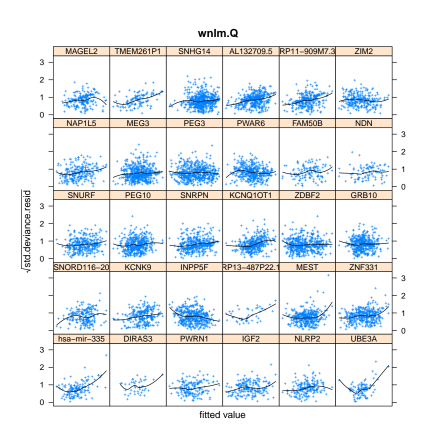


Figure S7: Checking the fit of wnlm. Q model: analysis of homoscedasticity.

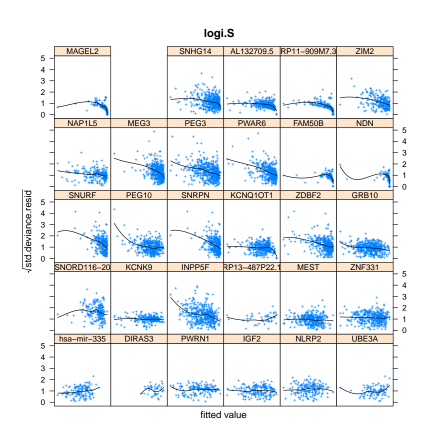


Figure S8: Checking the fit of logi.S model: analysis of homoscedasticity.

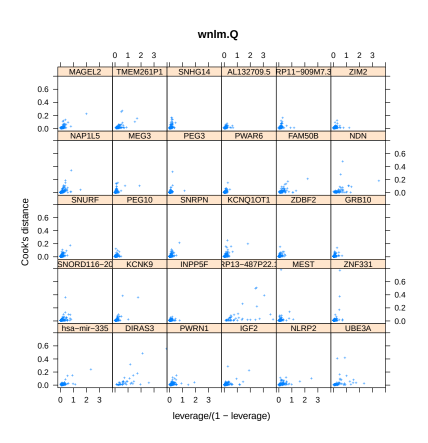


Figure S9: Checking the fit of wnlm.Q model: analysis of the impact of outliers.

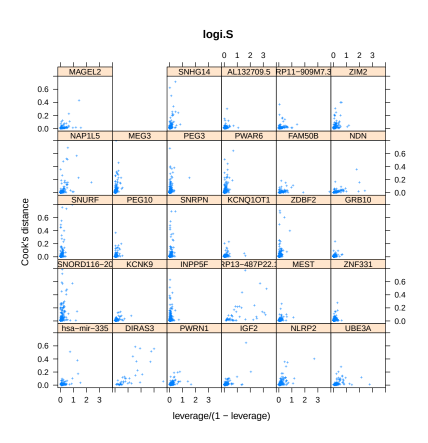


Figure S10: Checking the fit of logi.S model: analysis of the impact of outliers.

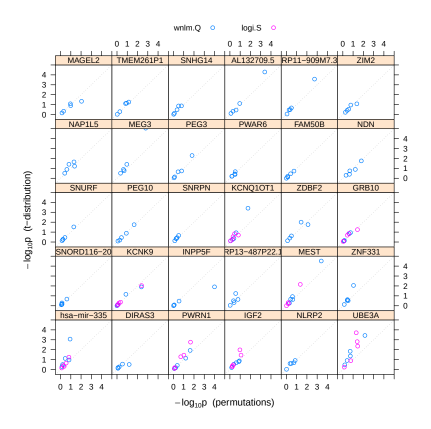


Figure S11: Agreement between a parametric (t-distribution) and non-parametric (permutations) method of estimating p-values.

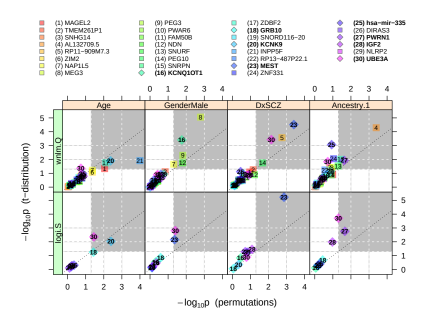


Figure S12: Significance of association between biological predictors and imprinted genes in the DLPFC calculated under wnlm.Q and logi.S. Under logi.S, only those genes are shown for which the model fit was acceptable.

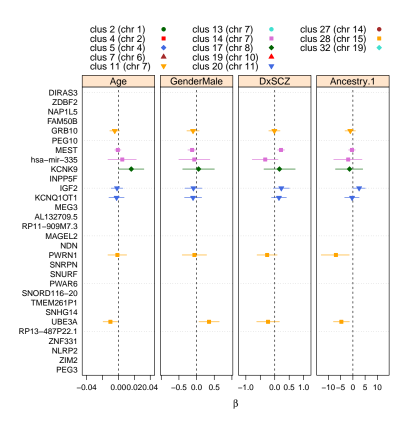
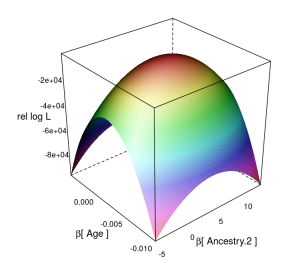


Figure S13: Estimate $\hat{\beta}_{jg}$ and 99% confidence interval of each regression coefficient β_{jg} under the logi.S model, where j is a predictor/term and g is a gene. $\hat{\beta}_{jg}$ and the confidence interval are shown only for those genes are shown for which the model fit was acceptable.



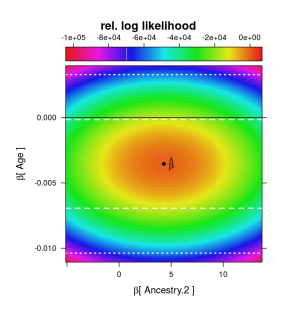


Figure S14: Top and bottom: two representations of the relative log likelihood on a 2 dimensional section of the p > 20 dimensional parameter space given the wnlm.Q model and data for the gene g = PEG3. The section was taken by fixing all but two parameters at their estimates: $\beta_{jg} = \hat{\beta}_{jg}$. A rectangular subspace for these two parameters, $\beta_{Age,g}$ and $\beta_{Ancestry,2,g}$, was chosen around the maximum likelihood estimate $\hat{\beta} = (\hat{\beta}_{Age,g}, \hat{\beta}_{Ancestry,2,g})$. The nearly parabolic shape of the log-likelihood function suggests that the regularity conditions for likelihood-based parametric inference are fulfilled.

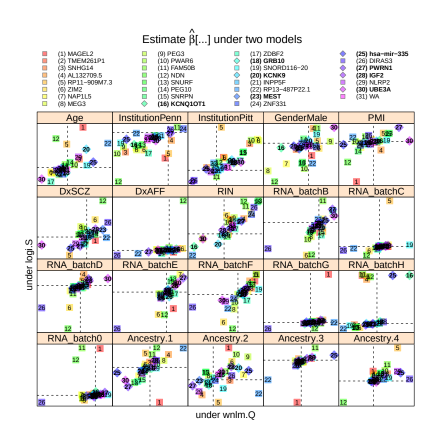


Figure S15: Agreement of the wnlm.Q and logi.S models in terms of estimated regression coefficients.

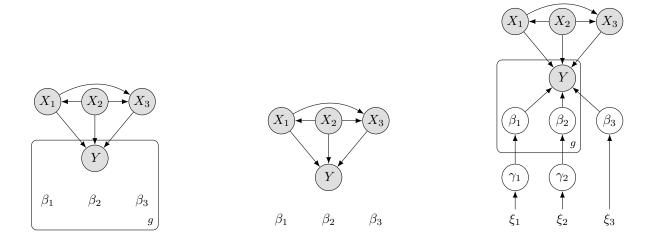


Figure S16: General dependency structure of three regression model frameworks. In all of threse model frameworks the regression coefficients $\beta_{1g},...,\beta_{3g}$ mediate, for a given gene g, probabilistic dependencies (arrows) between the response variable Y_g (read count ratio for g) and the corresponding predictors $X_1, ..., X_3$. For simplicity but without loss of generality only 3 predictors are depicted. The model frameworks differ in how $\beta_{jq_1}, \beta_{jq_2}, \dots$ relate to each other for a given predictor (or a given j). Left: there is no connection among $\beta_{jg_1}, \beta_{jg_2}, ...$ which means that the way Y_g , the read count ratio for gene g depends on predictor X_j is completely separate from how the read count ratio for any other gene g' (i.e. $Y_{g'}$) depends on it. Consequently no information may be shared among gene-specific models. Middle: In this case $\beta_{jg_1} = \beta_{jg_2} = ... \equiv \beta_j$ so that all genes are identical with respect to how their read count ratio depends the predictors. Thus genes share all information in the data in the sense that the model forces them to be identical. Right: Hierarchical Bayesian model where genes show both variation as well as invariance with regards to dependencies. The variation is described by the dependence of β_j on the hyperparameter γ_j , whereas the invariance by the dependence of γ_i on its hyperparameter ξ_i . Only this model framework allows information sharing among genes in a flexible way.

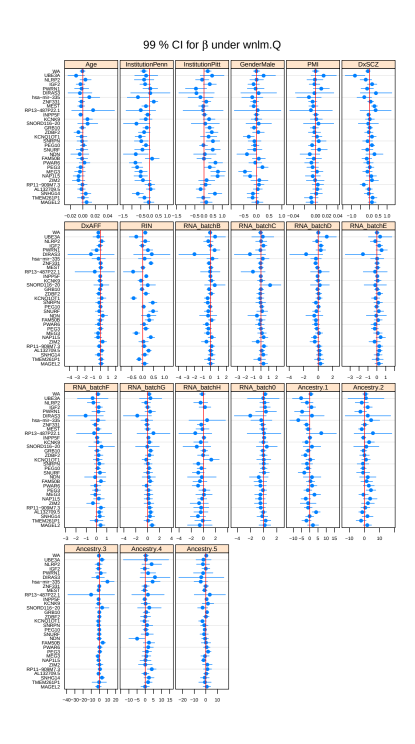


Figure S17: Estimates $\hat{\beta}_{jg}$ and confidence intervals for regression coefficients under the wnlm.Q model concerning for all predictors.

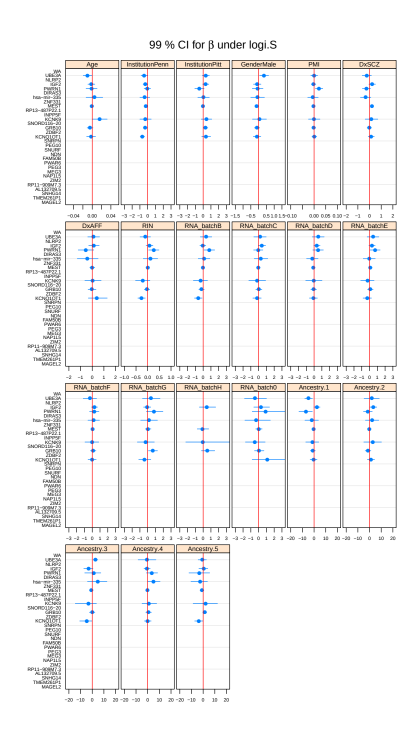


Figure S18: Estimates $\hat{\beta}_{jg}$ and confidence intervals for regression coefficients under the logi.S model concerning for all predictors. Gaps for certain genes indicate unacceptable fit.

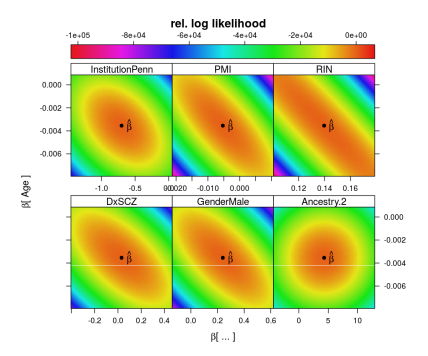


Figure S19: Analysis of orthogonality of regression coefficients. Relative log-likelihood surfaces for the gene PEG3 on various rectangles corresponding to 2D sections through the p > 20 dimentional parameter space. The set of points in a rectangle where log-likelihood takes the same value are quasi-ellipses, whose major and minor axes and their tiltedness express association between coefficients. For instance, β_{Age} is not associated with (orthogonal to) $\beta_{\text{Ancestry.2}}$ but is strongly associated with β_{RIN} . Such association hinders statistical inference due to the following circularity: If we knew the precise value of β_{RIN} we could estimate the true value of β_{Age} with higher precision and confidence but the precise value of β_{RIN} could only be obtained with high confidence if we precisely knew β_{Age} .

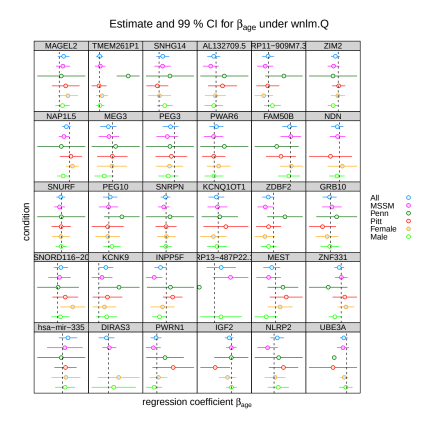


Figure S20: Analysis of interactions among predictors under the wnlm.Q model. Contextual dependence of the read count ratio on age, where the context is given by some specific level of Institution (MSSM, Penn, Pitt) or Gender (Female, Male).

Estimate and 99 % CI for β_{age} under logi.S

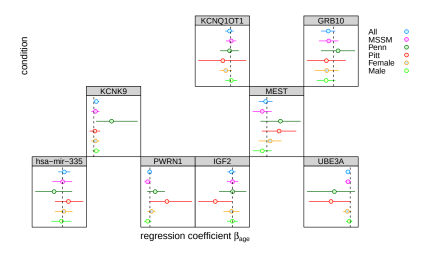


Figure S21: Analysis of interactions among predictors under the logi.S model. The missing panels correspond to genes for which logi.S did not provide acceptable fit.

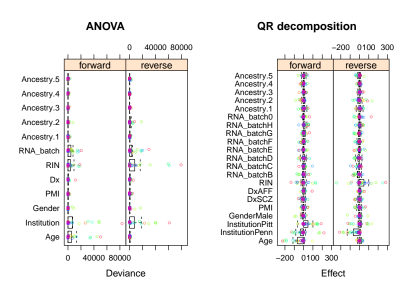


Figure S22: Left: analysis of variance (ANOVA) is undermined by the non-orthogonality of predictors because the reduction in deviance (i.e. in residual sum of squares) for a term (predictor) depends on the sequence in which terms are added to the model. Right: the same concept is demonstrated using QR decomposition.

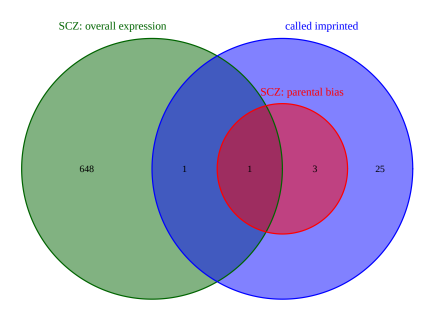


Figure S23: Association of genes' expression to schizophrenia (SCZ) assayed by two RNA-seq based approaches: total read count (overall expression, Nat Neurosci. 2016 Nov;19(11):1442-1453.) and read count ratio (allelic bias, present work). When these approaches are compared for only those genes that we find imprinted in the DLPFC in this study, 1 gene is found associated to schizophrenia by both approaches, 1 by only overall expression, and 3 by only allelic bias.