GWAS-based pathway analysis differentiates between fluid and crystallized intelligence

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ABSTRACT

Cognitive abilities vary among people. About 40-50% of this variability is due to general intelligence \((g)\), which reflects the positive correlation among individuals’ scores on diverse cognitive ability tests. \(g\) is positively correlated with many life outcomes, such as education, occupational status, and health, motivating the investigation of its underlying biology. In psychometric research, a distinction is made between general fluid intelligence \((g_F)\) - the ability to reason in novel situations - and general crystallized intelligence \((g_C)\) - the ability to apply acquired knowledge. This distinction is supported by developmental and cognitive neuroscience studies. Classical epidemiological studies and recent genome-wide association studies (GWASs) have established that these cognitive traits have a large genetic component. However, no robust genetic associations have been published thus far due largely to the known polygenic nature of these traits and insufficient sample sizes. Here, using two GWAS datasets, in which the polygenicity of \(g_F\) and \(g_C\) traits was previously confirmed, a gene- and pathway-based approach was undertaken with the aim of characterizing and differentiating their genetic architecture. Pathway analysis, using genes selected on the basis of relaxed criteria, revealed notable differences between these two traits. \(g_F\) appeared to be characterized by genes affecting the quantity and quality of neurons and therefore neuronal efficiency, whereas long term depression (LTD) seemed to underlie \(g_C\). Thus, this study supports the \(g_F\)-\(g_C\) distinction at the genetic level and identifies functional annotations and pathways worthy of further investigation.
INTRODUCTION

A large proportion (40-50%) of inter-individual variability in cognitive abilities is due to general intelligence \( (g) \), a quantitative trait that reflects the positive correlation among an individual’s scores on diverse cognitive ability tests (Deary, 2012; Spearman, 1904). A high \( g \) score is associated with many favorable life outcomes (Deary & Batty, 2011). Psychometric research distinguishes between general fluid intelligence \( (g_F) \) – the ability to reason in novel situations – and general crystallized intelligence \( (g_C) \) – the ability to apply acquired knowledge and learned skills (Carroll, 1993; Cattell, 1963). Although \( g_F \) and \( g_C \) are correlated at least 50% based on twin studies (Wainwright et al., 2005) and more so early and late in life (Li et al., 2004), developmental and cognitive neuroscience studies largely support the distinction between them. In normal aging, \( g_F \) declines earlier and more rapidly than \( g_C \) (Craik & Bialystok, 2006; Salthouse, 2004); in development, measures of verbal \( (g_C) \) and non-verbal \( (g_F) \) intelligence correlate differently with changes in brain structure (Ramsden et al., 2011). Furthermore, \( g_F \) is closely associated with fronto-parietal and anterior cingulate brain networks, while \( g_C \) is dependent on posterior frontal and temporal regions (Barbey et al., 2012; Glascher et al., 2009; Jung & Haier, 2007; Woolgar et al., 2010). Finally, at the population level, large gains in performance have been observed for tests that are strongly associated with \( g_F \), but not with \( g_C \) (Flynn, 2007).

More than half of the variability in intelligence tests is attributable to additive genetic effects (Deary et al., 2009a; Lee et al., 2010; Plomin & Spinath, 2004). In a recent genome-wide association study (GWAS), 40% and 51% of the phenotypic variability in \( g_C \) and \( g_F \), respectively, could be accounted for by genetic variants in linkage disequilibrium with common single nucleotide polymorphisms (SNPs) (Davies et al., 2011), providing a lower-
bound estimate of the narrow-sense heritability of these traits. Furthermore, using only SNP
data, ~1% of the variance in intelligence test scores in a sample could be predicted in an
independent sample (Davies et al., 2011). However, at the single marker level, no robust
genetic association with intelligence has yet been published, consistent with the observation
that the effect of individual SNPs may be too weak to be detected in complex polygenic traits
using the classical genome-wide $P \leq 5 \times 10^{-8}$ threshold (Wellcome Trust Case Control
Consortium, 2007) on the sample sizes currently available (International Schizophrenia
Consortium, 2009). Therefore, relaxing the significance criteria and exploiting the polygenic
signal by going beyond the traditional single-marker approach to gene- and pathway-based
methods may offer more power (Neale & Sham, 2004) and insight into the biological
processes underlying these traits (Wang et al., 2010).

The polygenic architecture of $g_F$ and $g_C$ was previously confirmed using the Cognitive Aging
Genetics in England and Scotland (CAGES) cohort as the discovery sample and the
Norwegian Cognitive NeuroGenetics (NCNG) adult lifespan sample for replication (Davies et
al., 2011). Here, using the same datasets, we combine single-marker, gene- and pathway-based
approaches to characterize the genetic architecture of $g_F$ and $g_C$ with respect to known
biological processes.

MATERIALS AND METHODS

GWASs: genotypes and phenotypes

The genotype and phenotype protocols and the samples have been described previously
(Davies et al., 2011; Espeseth et al., 2012). All participants gave written consent before the
study started. All procedures were conducted according to the tenets of the Declaration of
Helsinki and approved of by the relevant Research Ethics Committees. An overview of the genotype and phenotype protocols in the discovery and replication samples is provided below, with further details available in the original publications.

Discovery GWAS: The final NCNG GWAS consisted of 554,225 SNPs genotyped in a homogenous Norwegian sample of 670 individuals (457 females), ranging from 18 to 79 years of age (M = 47.6; SD = 18.3) (Espeseth et al., 2012). Participants completed a battery of psychometric tests, assessing general cognitive ability, memory, attention and speed of processing. The protocol was approved by the Regional Committee for Medical and Health Research Ethics, Southern Norway (project ID: S-03116). DNAs were genotyped on the Illumina Human610-Quad Beadchip. Quality control was performed with the “check marker” function of the R package GenABEL (Aulchenko et al., 2007). Individuals were excluded based on relatedness (“ibs.threshold”=0.85), heterogeneity, unresolved sex discrepancies and call rate ≤ 0.97. Population structure was assessed by multidimensional scaling (MDS) analysis (using 100,000 random SNPs), removing outlying samples with possible recent non-Norwegian ancestry. No additional adjustment for population structure was performed due to the homogeneity of the sample (Espeseth et al., 2012). SNPs with a call rate ≤ 0.95, minor allele frequency ≤ 0.01 and Hardy-Weinberg Equilibrium (exact test) P-value ≤ 0.001 were excluded.

Replication GWAS: The CAGES sample consists of five cohorts, the Lothian Birth Cohorts of 1921 (Deary et al., 2009b; Deary et al., 2004) and 1936 (Deary et al., 2007) (LBC1921, LBC1936), the Aberdeen Birth Cohort of 1936 (ABC1936) (Deary et al., 2009b; Deary et al., 2004) and the Manchester and Newcastle Longitudinal Studies of Cognitive Aging cohorts (Rabbitt et al., 2004). The final GWAS dataset consisted of 549,692 SNPs genotyped in 3511
healthy individuals (2115 females) with detailed cognitive ability measurements taken in middle to older adulthood (mean age ranged from 64.6 in the ABC1936 to 79.1 in LBC1921; overall age range: 44-93) (Davies et al., 2011). Ethical approval was obtained from the relevant Research Ethics Committees. Individuals were excluded based on unresolved gender discrepancy, coefficient of relatedness>0.025 (based on 549,692 autosomal SNPs), call rate≤0.95 and evidence of non-Caucasian descent as determined by MDS analysis (Davies et al., 2011). SNPs were included if they met the following conditions: call rate≥0.98, minor allele frequency≥0.01 and Hardy–Weinberg equilibrium test with P≥0.001. Population structure was assessed by MDS analysis, and four MDS components were fitted as covariates to correct for any population stratification that might be present.

Cognitive phenotypes: Different measures of gF and gC were employed for each of the five CAGES samples and the NCNG sample (Davies et al., 2011). In general, the different tests of gF aimed to assess each individual’s capacity to reason logically and solve problems in novel situations, relatively independently of acquired knowledge. gF employs aspects of processing speed, attention, memory and executive function. The gC tests were vocabulary-based, assessing each individual’s semantic knowledge. All measures were corrected for sex (with the exception of the CAGES Manchester and Newcastle gF, which was derived separately for males and females) and age. The standardized residuals were then extracted and used as the trait measures in all subsequent analyses. In the NCNG, of the 670 individuals, 629 and 643 had scores for gF and gC, respectively. In the CAGES, of the 3511 individuals, 3400 and 3482 had scores for gF and gC, respectively.

Analyses

Construction of gene lists
A ‘bottom-up’ approach (Liu et al., 2007) was undertaken. Both single-marker and gene-based association analyses were performed, testing SNPs and genes individually first in order to construct a list of genes that would then be subjected to pathway analysis, via IPA, to identify over-represented functions and/or pathways. For each of gF and gC, genetic factors that showed evidence of association in the NCNG were identified and filtered further based on evidence of replication in the CAGES. Single-marker and gene-based statistical methods and thresholds were selected and applied, as appropriate, with the aim of controlling the Type II (false-negative) as well as the inversely related Type I (false-positive) error rates, incorporating instead prior biological knowledge to the interpretation of the findings and generating testable hypotheses for further investigation (Lieberman & Cunningham, 2009; Williams & Haines, 2011).

The LDsnpR tool (Christoforou et al., 2012a) was used to (1) annotate individual SNPs and (2) assign SNPs to genes for the gene-based analyses. SNPs were assigned to genes (Ensembl 54 definitions) if they were located physically within the boundaries of the gene (+/-10kb) or if they were in high linkage disequilibrium (LD; r²≥0.80 based on HapMap CEU (http://hapmap.ncbi.nlm.nih.gov/)) with another SNP located within the boundaries of the gene (+/-10kb).

Single-marker analyses: In the NCNG sample, association between individual SNP alleles and cognitive phenotypes was tested using linear regression analysis (1 d.f. coefficient t-test), as implemented in PLINK (Purcell et al., 2007). The CAGES GWAS data comprised the inverse variance weighted model ‘meta P-values’ produced in the original CAGES five-sample meta-analysis (i.e. as reported by Davies et al., 2011). A meta-analysis of the CAGES
and NCNG single-marker allele P-values was performed on the overlapping SNPs, using an inverse variance weighted model, as implemented in METAL (Willer et al., 2010).

All SNPs with an asymptotic nominal P-value ≤0.05 in the NCNG were then mined for replication in the CAGES GWAS data. As advised in Konig (2011), positive single-marker replication was determined on the basis of P≤0.05 and the same direction of effect in both the NCNG and CAGES samples, resulting in a meta-analysis P-value that is more significant than either of the original P-values. Using LDsnpR, all SNPs that replicated were annotated, where possible, with a gene name or ENSEMBL identifier and taken forward for pathway-based analysis by IPA.

**Gene-based analyses:** All 554,225 SNPs in the NCNG GWAS were assigned, where possible, to genes using LDsnpR. For each gene containing a SNP, association was assessed by applying PLINK’s permutation-based set test on the LDsnpR-generated sets. Gene-based statistics generated by permutation-based methods automatically account for potential confounding factors, such as LD structure and gene length or SNP number, through the generation of an empirical null distribution (Liu et al., 2010). Since the true underlying genetic architecture of these traits is unknown, three different models were tested to obtain a gene-based association score (Lehne et al., 2011):

1. the minimum P-value (minP) model, which assigns to each gene the association statistic, or P-value, of the most significant SNP in the gene, assumes that a single SNP within the gene contributes to the phenotype;

2. the all P-values model, which assigns to each gene the mean association statistic of all the SNPs in the gene, assumes that all or most SNPs within the gene contribute to the phenotype;
(3) the $P \leq 0.05$ threshold model, which assigns to each gene the mean association statistic of all SNPs within the gene that have a $P \leq 0.05$, assumes that only a few or a subset of SNP contribute to the phenotype.

The $P$-values were computed based on 10,000 permutations, and the lowest of the three permuted $P$-values was retained as the gene-based $P$-value for each gene. No additional correction was applied to the gene-based score to account for testing three models due to the high correlation between the tests. Genes with gene-based permuted $P \leq 0.05$ were mined for replication in the CAGES.

In order to preserve the Type II error rate, the replication criteria in the CAGES were relaxed to include any gene that contained at least one SNP with a ‘meta-$P$’ $\leq 0.05$, ensuring that significance under any genetic architecture was captured. In the NCNG, for example, all genes with a minimum gene-based permuted $P \leq 0.05$ contained at least one SNP with a $P \leq 0.05$ (data not shown). This replication criterion of at least one SNP with a ‘meta-$P$’ $\leq 0.05$ is equivalent to the unadjusted minimum $P$-value approach. Thus, the CAGES SNPs were first assigned to genes using LDsnpR, as described above, and the genes were scored using the minimum $P$-value approach, assigning to the gene the $P$-value of the most significant SNP in that gene (without further adjustment). All genes that were selected from the NCNG and that also replicated in the CAGES (i.e. meta-$P \leq 0.05$) were taken forward for analysis with IPA.

Finally, after the relevant gene lists were pruned for LD on the basis of their association signals and position, the hypergeometric distribution, as implemented in www.geneprof.org/GeneProf/tools/hypergeometric.jsp, was used to assess whether the number of genes replicating in the CAGES was greater than expected by chance.
**Construction of gene lists – stringent replication criteria:** Since the aforementioned single-marker and gene-based replication criteria in the CAGES are prone to inflating the Type I error rate due to gene-length bias, the robustness of the subsequent main IPA findings was assessed by re-running IPA on gene lists constructed using more stringent statistical thresholds and methods. Firstly, for the single-marker analysis, only genes tagged by the SNPs that met the previously described replication criteria and resulted in a meta-analysis \( P \leq 1 \times 10^{-5} \), as supported by the National Human Genome Research Institute GWAS catalog (Welter et al., 2014), were taken forward for IPA analysis. For the gene-based analysis, replication in the CAGES was determined using two gene-scoring methods which accounted for the number of SNPs per gene and, critically, for LD without requiring genotype-level data. The first method scored each gene with the most significant P-value (i.e. the minP approach), adjusting for the number of SNPs using a modified Sidak’s correction (Saccone et al., 2007). This approach has been shown to perform as well as regression-based methods (Segre et al., 2010; Christoforou et al., 2012a). The second approach was comparable to the regression-based ‘all P-values’ model described above, combining all SNPs mapped to the gene using Brown’s approximation method, as implemented in PLINK (Moskvina et al., 2011; Purcell et al., 2007). The NCNG genotype data was used to estimate the LD between SNPs. Thus, the genes with gene-based permuted \( P \leq 0.05 \) in the NCNG which met a \( P \leq 0.05 \) with either of these two methods in the CAGES were subjected to IPA analysis.

**Ingenuity Pathway Analysis (IPA)**

Genes that showed evidence of association in the NCNG and of replication in the CAGES based on either the single-marker and/or the gene-based analysis were selected for pathway analysis with IPA (Ingenuity® Systems, www.ingenuity.com) to identify the most relevant biological functions and pathways. IPA gathers information from published data from ~3600
peer-reviewed journals regarding relationships between genes and proteins. The information is updated weekly, manually curated and stored within the Ingenuity® Knowledge Base, which is then queried during an analysis to identify specific biological functions, or “function annotations”, and pathways enriched within the submitted gene list.

In this study, the Ensembl 54 gene identifiers were uploaded into IPA and mapped, if possible, to their corresponding object in the Ingenuity® Knowledge Base (Genes Only). A “Core Analysis” was performed, including both direct and indirect relationships and using all available data sources in all species available. Molecules and/or relationships that were either experimentally observed or predicted with high confidence were considered. For the main analyses, which aimed to differentiate gF and gC as it relates, most relevantly, to the nervous system, tissues and primary cells were restricted to those of the nervous system and cell lines were restricted to those of the central nervous system. Additional analyses were performed to evaluate the robustness and relevance of the main findings, including (1) re-running the IPA having excluded genes known to be extensively studied to assess the possibility of publication bias driving the main findings and (2) re-running the IPA having included all tissues and cell lines, except those specific to cancer, to assess the relevance of the main findings in the context of more global annotation.

In particular, we focused on the significant function annotations. The function annotations are organized based on Ingenuity® Ontology, which consists of a manually built and maintained hierarchical data structure of hundreds of thousands of unique classes. As part of the ‘Core Analysis’, IPA performs a Fisher Exact Test (FET) to determine whether the submitted gene list consists of genes of a particular function annotation more than expected by chance given the proportion of genes of that particular function annotation in the entire Ingenuity
Knowledge Base. It also uses the Benjamini-Hochberg (BH) multiple testing correction method to adjust for the number of ontologies tested, providing a false discovery rate for a particular function annotation. In this study, all functional annotations with a FET \( P \leq 0.05 \) (and corresponding BH \( P \leq 0.25 \)) were considered significant.

**RESULTS**

**Construction of gene lists**

*Single-marker analyses*: A total of 554,225 SNPs were tested for allelic association to gF (N=629 individuals) and gC (N=643 individuals) in the NCNG sample. As these GWAS data results were not presented in the previous study (Davies *et al.*, 2011), the traditional SNP-level diagnostics and results are provided in the Supporting Information (Figure S1 and Dataset S1). The genomic inflation factor was 1 for both gF and gC (Figure S1), indicating that the data did not suffer from population stratification or other systematic bias. When mining the CAGES data, 816 SNPs in gF and 884 SNPs in gC met the replication criteria of \( P \leq 0.05 \) and the same direction of effect (Supporting Information Dataset S1). Using the LD-based binning approach implemented in LDsnpR (Christoforou *et al.*, 2012a), 481 (59%) of the gF SNPs and 549 (62%) of the gC SNPs were assigned to Ensembl 54 genes. 159 gF SNPs and 160 gC SNPs mapped to more than one gene on the basis of LD. Since it is not possible to identify the true source of the association on the basis of the SNP P-value alone (Christoforou *et al.*, 2012a,b), all genes were retained for pathway analysis. The significant function annotations or pathways were subsequently manually examined to ensure that they did not consist of clusters of genes representing the same genetic association signal. Thus, 503 and 530 Ensembl 54 genes for gF and gC, respectively, (Supporting Information Dataset S1) were taken forward for IPA analysis. 171 (34%) of the gF genes and 212 (40%) of the gC genes were implicated by more than one replicated SNP.
Gene-based analyses: A gene-based approach was also used to identify candidate genes for pathway analysis, allowing for locus heterogeneity and the aggregation of multiple weaker association signals. In the NCNG, of the 34,109 eligible Ensembl 54 gene entries, 2698 and 2615 met the nominal significance threshold of set-based permuted $P \leq 0.05$ in $gF$ and $gC$, respectively (Supporting Information Dataset S2). Of these, 841 $gF$ genes and 920 $gC$ genes had a minimum $P$-value gene-based score of $P \leq 0.05$ in the CAGES (Supporting Information Dataset S2). After the relevant gene lists were first pruned for LD based on position and association signal, resulting in 607 and 652 relatively independent association signals for $gF$ and $gC$, respectively, the number of genes showing evidence of replication was more than expected by chance for $gC$ (hypergeometric $P=4.5 \times 10^{-4}$), but not for $gF$ (hypergeometric $P=0.25$). The 841 $gF$ genes and 920 $gC$ genes were added to the list of genes identified by single-marker analysis and taken forward for IPA.

Ingenuity Pathway Analysis

Figure 1 summarizes the number of SNPs and genes identified in each analysis and the relevant corresponding overlaps. 1182 genes for $gF$ and 1294 genes for $gC$ were identified through both the single-marker and gene-based analyses and subjected to IPA analysis. Of these genes, 853 for $gF$ and 893 for $gC$ were identified by their Ensembl Gene identifier in the IPA database (i.e. “IPA Ready” genes; Supporting Information Dataset S3) and were thus available for pathway analysis. 117 genes were in common between the two traits.

IPA’s “Core Analysis” was run on the two lists of genes, restricting the analysis to tissues and cell lines related to the nervous system. Function annotations that were significantly enriched with a Fisher Exact Test (FET) $P \leq 0.05$ and a Benjamini-Hochberg (BH) $P \leq 0.25$, which
account for all of the function annotations tested, were declared significant and evaluated. For both traits, function annotations involved in the biological function “Nervous System Development and Function” were the most common, accounting for >25% of the function annotations. These were followed by annotations involved in “Cell-to-Cell signaling” and “Cellular Assembly and Organization” in both gF and gC (~8%) (Supporting Information Dataset S3). However, it is important to note that some function annotations are categorized under multiple biological functions, resulting in redundancy in the data (e.g., “axonogenesis” appears under “Nervous System Development and Function”, “Cell Morphology”, “Cellular Function and Maintenance”, “Cellular Assembly and Organization” and “Tissue Development”). When considering only the non-redundant function annotations, specifically those identified on the basis of at least two genes from the inputted gene list, 85 and 54 function annotations were identified as significantly enriched in gF and gC, respectively (Supporting Information Dataset S3).

Only four function annotations were enriched in both gF and gC, including “microtubule dynamics” and “chemotaxis of neurons”, which could be generally categorized as architectural features of neuronal development. The set of genes leading to the enrichment of these overlapping annotations in gF was different to that in gC (Table 1), except for the “migration of GABAergic neurons” annotation which was attributed to the same two genes in gF and gC.

The most statistically significant functional annotations were identified in gC. “Synaptic depression” ranked at the top (FET P=2.9x10^{-6}; BH P=0.0015), with the related “long term depression” in general (LTD; FET P=2.0x10^{-5}; BH P=0.0052) and LTD of specific cells (Table 2), all of which are highly related as indicated by the composite genes. “Guidance of
“axons” and “schizophrenia” were also significantly enriched in gC (FET P=4.8x10^{-4} and 0.042, respectively), the former also ranking third (Table 2 and Supporting Information Dataset S3).

The general functional enrichment profiles of gF and gC were different. No function emerged as notably significant in gF, a finding which was further emphasized by the ‘flat’ multiple-testing corrected BH P-value of 0.18 for all significant annotations. The most significantly enriched functional annotation identified was “synaptic fatigue of synapse” (FET P=0.0021), followed by “apoptosis of spinal cord cells” (FET P=0.0041) and “inhibition of neurons” (FET P=0.0053) (Table 3). However, the predominance of function annotations relating to “quantity”, particularly of neurons and of other structures of the nervous system was notable (Table 3, Supporting Information Figure S2 and Dataset S3). Other functions were related to the quantity (e.g., formation, loss, survival, and apoptosis), quality (e.g. synaptic fatigue, degradation, atrophy and myelination), or morphology of neurons or related structures. Long-term potentiation (LTP) was another repeated function for gF (Supporting Information Figure S2). One gene, brain-derived neurotrophic factor (BDNF), which showed gene-based association with gF (Supporting Information Dataset S2), contributed to the enrichment of several significant function annotations for gF (Table 3 and Supporting Information Dataset S3). As BDNF is one of the most extensively studied genes in the field (Green et al., 2008), it could bias the IPA results, which are based on peer-reviewed publications. However, when IPA was run without BDNF, most of the significant findings withstood its exclusion (Table 3 and Supporting Information Dataset S3) and the general functional enrichment profile observed in gF was preserved. A manual examination of the chromosomal positions of the sets of genes leading to the enrichment of the significant annotations listed in Tables 1-3
ensured that these significant enrichments are due to independent association signals and not to clusters of genes in LD representing the same association signal.

In order to gauge the robustness of the above findings in the absence of the gene-length bias that may have been introduced with the relaxed replication criteria, the CNS-specific IPA analysis was also performed on gene lists constructed on the basis of more stringent replication criteria. For gF, the single-marker analysis identified two genes which contained replicated SNPs that met the meta-analysis $P \leq 1 \times 10^{-5}$ threshold (Supporting Information Dataset S1). The gene-based analysis identified 178 genes with gene-based $P \leq 0.05$ using either the modified Sidak approach or Brown’s approximation method (Supporting Information Dataset S2). Together, this resulted in a total of 180 genes available for IPA analysis for gF. For gC, 5 and 224 genes were identified via the single-marker and gene-based analyses, respectively, resulting in 225 unique genes available for IPA analysis. Nine genes were common to both gF and gC.

The IPA results emerging from the abridged gene lists were not as impressive in terms of the number of significant unique function annotations ($N=26$ and $8$ for gF and gC, respectively) and in terms of their general enrichment profiles for gF and gC (Supporting Information Dataset 3). Also, for both traits, different function annotations emerged as most significant, namely “neuritogenesis” for gF (FET $P=2.8 \times 10^{-4}$) and “hypoplasia of cerebellar vermis” (FET $P=0.0015$) (Supporting Information Dataset 3). However, the main functions which were found to distinguish gF from gC in the original, less conservative analysis remained significant. “LTD” and “synaptic depression” remained unique to gC albeit at a reduced significance level (FET $P=0.018$ and $0.047$, respectively) (Table 2 and Supporting Information Dataset 3). For gF, functions which related to quantity, quality or morphology of
neurons and synapses still predominated. These included “quantity”, “morphogenesis” and “development” of neurons, synapses and other brain structures (Supporting Information Dataset 3). No function annotations were common to both traits in this analysis.

Finally, returning to the original gene lists based on the relaxed criteria, IPA was run again on both traits, using information from all cell lines, with the exception of cancer cell lines, in the Ingenuity® Knowledge Base. This helped to evaluate the primary CNS-restricted results relative to more global annotation, to identify annotations that may be related to cognitive ability and to enable the inclusion of published studies of brain-related traits that were executed in non-CNS cell lines, such as lymphoblastoid cell lines (Gladkevich et al., 2004). The significance of the original, CNS-restricted findings was largely unaffected. The most significant annotations for gF in this analysis fell under the Disease and Disorder Category of Cardiovascular Disease (e.g. “vascular disease”, FET \( P = 7.1 \times 10^{-6} \); Supporting Information Dataset S3). In addition, the function annotations “development of brain”, “behavior”, “cognition disorders”, “cognitive impairment” and “schizophrenia” were also significantly enriched in gF (Supporting Information Dataset S3). For gC, “synaptic depression” remained the most significant annotation. However, it was followed by the Cardiovascular Disease function annotation “coronary artery disease” (FET \( P = 1.8 \times 10^{-6} \); Supporting Information Dataset S3). The significance of the enrichment of “schizophrenia” also marginally improved (from FET \( P = 0.042 \) to FET \( P = 0.026 \), Supporting Information Dataset S3).
DISCUSSION

Novel polygenic approaches to analysing GWAS data have greatly improved our understanding of complex traits and have captured more of the “hidden heritability” (Davies et al., 2011; Visscher et al., 2012; Maher, 2008; Manolio et al., 2009) but have failed to identify the markers or genes implicated. Pathway-based techniques are complementary polygenic methods that support biological analysis of GWAS data. Gene-based methods, which are ideal for pathway-based approaches (Liu et al., 2010), permit the aggregation of SNPs of smaller effect and test the gene as a whole, potentially increasing power (Neale & Sham, 2004). These analyses are also more permissive to locus heterogeneity, where multiple independent variants within a locus have independent effects on a trait (Christoforou et al., 2012a; Yang et al., 2012), and they use prior biological knowledge, facilitating a more meaningful interpretation of data (Wang et al., 2010). Therefore, if the genetic signals underlying gF and gC, two highly heritable and polygenic traits, cluster in known biological pathways, it should be possible to use pathway-based analyses to identify which biological processes are most strongly implicated in these subcomponents of g. We particularly wanted to determine whether gF and gC could be differentiated at the level of biological pathways or functions as they have been shown to be distinct in terms of development and brain structure. Taking a ‘bottom-up’ approach (Liu et al., 2007), we analysed two GWAS datasets using both single-marker and gene-based analyses to construct gene lists for IPA analysis. Although some significantly enriched functions overlapped in the two traits, the overall picture suggested distinct functional enrichment profiles, supporting the gF-gC distinction at the genetic level.

The most statistically significant finding was the enrichment in gC of genes involved in synaptic depression and LTD. This finding, which was unique to gC, also survived the FDR
multiple testing correction at the more conservative 0.05 threshold and the application of more stringent replication criteria for the construction of gene lists. It was also the top finding when the pathway analysis included non-CNS-related tissue types and conditions. LTD and long-term potentiation are the major forms of long-lasting synaptic change in the mammalian brain (Collingridge et al., 2010). LTD is involved in synaptic pruning during development (Peineau et al., 2007) and is thus important in adult neuroplasticity. Synaptic LTD is mediated by the effect of L-glutamate and other neurotransmitters on several types of receptors. The resulting synaptic plasticity is necessary for hippocampus-dependent learning and memory, certain types of behavioral flexibility and novelty detection (Collingridge et al., 2010). LTD deficits have been associated with reduced working memory and reversal of memory performance in rats. Studies focusing on medial temporal lobe regions have revealed a role of LTD in memory stabilization (in the amygdala) (Migues et al., 2010) and recognition memory (in the perirhinal cortex) (Winters & Bussey, 2005), implicating LTD in the development and maintenance of knowledge representations.

For gF, the association signal was less striking in terms of statistical significance. gF was predominantly characterized by genes that control the quantity, morphology and integrity of neurons and synapses. These factors affect the quality and efficiency of neuronal signaling (Brown et al., 2008), which, in turn, affect cognition, as indicated by reduced activation in individuals with higher cognitive ability (Prat et al., 2007; Reichle et al., 2000). Since gF declines in cognitive ageing, the enrichment of these functions is consistent with the decline in synapse number, brain volume, and white matter integrity in the ageing brain (Fjell et al., 2009; Morrison & Hof, 1997; Walhovd et al., 2011; Westlye et al., 2010).
Only four function annotations were common to both traits. These were related to development and structural aspects (e.g., microtubule dynamics, dendrite formation), which play central roles in synapse formation and are thus likely to be important for cognitive function (Bramham et al., 2010). The genes leading to enrichment of three of these functions were different for gF and gC, reflecting the gain of information that gene-set or pathway-based approaches offer by enabling the identification of overlap between related traits or replication of the same trait.

The function annotation “schizophrenia” was also significantly enriched in gF and gC, indicating that the identified genes have also been implicated in schizophrenia by other studies. A polygenic risk score for schizophrenia was recently shown to be associated with lower IQ at age 70 and greater decline in IQ level in one of the CAGES cohorts (Mcintosh et al., 2013), confirming previous observations at the single candidate level of a genetic relation between general cognition and schizophrenia (Toulopoulou et al., 2010). Furthermore, enrichment of the terms “behavior”, “cognition disorders” and “cognitive impairment” suggests that the genes identified here have been implicated in other studies of cognition. Finally, annotations relating to “cardiovascular disease” ranked in the top two in both gF and gC. Cognitive dysfunction is well documented in patients with cardiovascular disease (Vogels et al., 2007).

The heritability for gF and gC are similar, but the strength of association, extent of replication and strength of enrichment were all greater for gC than for gF (Davies et al., 2011). There are several possible reasons for this. First, gF may be more heterogeneous at the phenotype level. While gC was defined as the standardized score of a single measure in each sample, gF estimation was based on a hierarchical principal component analysis from an array of specific
measures. The idea that the psychometric structure of gF is more heterogeneous is consistent with another major model of intelligence (Johnson & Bouchard, 2005; Vernon, 1964), in which verbal abilities are retained as a single second stratum component, while fluid abilities are separated into perceptual and mental rotation skills, thus reflecting higher phenotypic complexity. Second, assessment of gF was not based on identical subsets of specific tests in NCNG and each of the CAGES subsamples. However, the different batteries of cognitive tests yield almost identical estimates of general intelligence (Johnson et al., 2004). Third, the age differences between the samples could have a differential effect given that gF decreases with age while gC is relatively stable, even though the correlation between them increases with age (Li et al., 2004). Finally, the difference in enrichment profiles may be magnified by gene-length bias, given that a stronger association was identified for gF when IPA was re-run using the genes that passed the more stringent criteria.

The main complicating factor in this and other studies of brain-related traits is gene-length bias, as it presents an analytical ‘Catch-22’. While it is important to correct genes for their length, or equivalently, the number of SNPs tested, to control for false findings, doing so risks over-penalizing and thus eliminating the most relevant candidate genes and therefore pathways for intelligence. It is recognized that gene length is related to functional complexity (Xu et al., 2006) and it is known that brain-expressed genes involved in relevant neuronal processes and/or associated with autism and intellectual disability are substantially longer (King et al., 2013; Michaelson et al., 2012). Also, it has been suggested that longer genes are larger targets and therefore more prone to random mutation and are subject to different transcriptional mechanisms that may be functionally relevant to brain-related traits (King et al., 2013; Solier et al., 2013). Thus, in order to balance the competing Type I and II error rates, we focused our analysis on a well-accepted and recommended discovery-replication
approach (Jia et al., 2011), running the replication in one of the largest samples of its kind (the CAGES). Strict replication criteria were used in the single-marker analyses (Konig, 2011). For the gene-based analyses, a robust permutation-based approach, which accounts for LD structure and SNP number, was used to select genes for replication, the criteria for which were subsequently relaxed to avoid over-penalizing the larger, more relevant genes. In order to then assess the robustness of these findings, IPA was also run on genes that survived the more stringent approach which corrects for SNP number. Since the power of the IPA analysis was greatly compromised by the ~6-fold reduction in the number of genes available for the analysis, it was not surprising that the evidence for the genetic distinctiveness of the two traits was weaker. Nevertheless, the main distinguishing features between gF and gC were upheld. This suggests that the true genetic architecture of these two traits may lie in the middle, and is inaccessible using current approaches which either ignore or over-penalize for the length of the gene.

As larger consortia form and pathway analyses continue to improve, a better understanding of the genetic architecture of gF and gC will emerge (Lencz et al., 2014; Khatri et al., 2012). Meanwhile, our study serves as a starting point, supporting the gF-gC distinction at the genetic level and critically converging with the findings of developmental and cognitive neuroscience studies. The specific function annotations, or pathways, identified are worthy of further replication and interrogation, using, for example, ‘top-down’ approaches such as gene set enrichment analysis (Fernandes et al., 2013) to test specific gene sets constructed on the basis of these findings.
REFERENCES


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Wellcome Trust Case Control Consortium (2007) Genome-wide association study of 14,000 cases of seven common diseases and 3,000 shared controls. Nature 447, 661-678.


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FIGURE LEGEND

Figure 1. Summary of SNP- and gene-based analyses and corresponding numbers of genes identified for IPA for both $gF$ (left) and $gC$ (right). The boxed numbers in the centre represent the number of genes identified as a result of the SNP-based analyses (from top, down), the gene-based analyses (from bottom, up), the resulting total number of unique genes submitted to IPA and the number of relevant overlapping genes (in parentheses). * indicates that the GWAS SNPs were assigned to genes by LDsnpR.
Table 1. Significant IPA Function Annotations in both $g_F$ (left) and $g_C$ (right).

<table>
<thead>
<tr>
<th>Function</th>
<th>Function Annotation (FA)</th>
<th>$g_F$ FET P-value</th>
<th>$g_F$ BH P-value</th>
<th>Genes</th>
<th>$g_C$ FET P-value</th>
<th>$g_C$ BH P-value</th>
<th>Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>chemotaxis</td>
<td>chemotaxis of neurons</td>
<td>0.025</td>
<td>0.18</td>
<td>BDNF, GFRA1, RGS3</td>
<td>0.0037</td>
<td>0.17</td>
<td>EPHB2, GFRA1, SEMA3A, SLIT2</td>
</tr>
<tr>
<td>formation</td>
<td>formation of dendrites</td>
<td>0.028</td>
<td>0.18</td>
<td>ACHE, BCL11A, BDNF, CTNNA2, GRIN3A, NRG1, PRKG1, SGK1, SYNE1</td>
<td>0.035</td>
<td>0.22</td>
<td>DAB1, DSCAM, KLF7, KNDC1, MAP2, NRG1, PPP1R9B, RAC2, SEMA3A</td>
</tr>
<tr>
<td>organization</td>
<td>microtubule dynamics</td>
<td>0.030</td>
<td>0.18</td>
<td>ACHE, ATXN10, BAX, BCL11A, BCL2, BDNF, BSN, CDH1, CNTN4, CNTNAP2, CTNNA2, CTNNB1, CYP19A1, DISC1, EGFR, GDA, GRIN3A, GSN, IGF1R, KATNB1, MAPK8, NFIB, NRG1, PRKG1, PTPRM, RND1, SGK1, SLC18A3, SLIT1, SLIT3, SNCA, SYNE1, UHMK1</td>
<td>0.032</td>
<td>0.22</td>
<td>ATXN2, BBS10, CDH4, CNTN4, DAB1, DCC, DSCAM, EPHB1, EPHB2, GPM6A, KLF7, KNDC1, LAMB1, LRRC7, MAP2, MBP, MTOR, NRG1, PCDH15, PLD1, POU3F1, PPP1R9B, PRKCE1, PRKCA, PTPRM, RAC2, RIT2, RUFY3, SEMA3A, SLIT1, SLIT2, TNFRSF12A, TRPV4, VANG</td>
</tr>
<tr>
<td>migration</td>
<td>migration of GABAergic neurons</td>
<td>0.012</td>
<td>0.18</td>
<td>GFRA1, SLIT1</td>
<td>0.013</td>
<td>0.22</td>
<td>GFRA1, SLIT1</td>
</tr>
</tbody>
</table>

For each general function, the specific function annotation (FA) is shown together with the corresponding Fisher Exact Test (FET) P-value, the Benjamini-Hochberg (BH) multiple testing corrected P-value and the genes responsible for the enrichment signal in the function annotation.

Individual genes that contribute to the enrichment of a particular FA in both $g_F$ and $g_C$ are in bold.
Table 2. Top 25 IPA Function Annotations for gC

<table>
<thead>
<tr>
<th>Function</th>
<th>Function Annotation (FA)</th>
<th>FET P-value</th>
<th>BH P-value</th>
<th># Genes</th>
<th>Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>synaptic depression</td>
<td>synaptic depression</td>
<td>2.9x10^-7</td>
<td>0.0015</td>
<td>13</td>
<td>ADCY5, ADCY8, CNR1, DRD5, EPHB1, EPHB2, GRM7, LRRC7, MTOR, PRKCA, RYR3, ST8SIA4, SYNJ1, (ITSN1)</td>
</tr>
<tr>
<td>long term depression</td>
<td>long term depression</td>
<td>2.0x10^-5</td>
<td>0.0052</td>
<td>11</td>
<td>ADCY5, ADCY8, CNR1, DRD5, EPHB2, GRM7, LRRC7, MTOR, PRKCA, RYR3, ST8SIA4</td>
</tr>
<tr>
<td>guidance</td>
<td>guidance of axons</td>
<td>4.8x10^-3</td>
<td>0.061</td>
<td>16</td>
<td>ALCAM, ANK3, CDH4, CNTN4, DCC, EPHB1, EPHB2, ERBB4, EXT1, GLI3, KLF7, NFASC, PLXNA2, SEMA3A, SLIT1, SLIT2</td>
</tr>
<tr>
<td>long term depression</td>
<td>long term depression of cells</td>
<td>4.8x10^-3</td>
<td>0.061</td>
<td>6</td>
<td>ADCY5, CNR1, DRD5, PRKCA, RYR3, ST8SIA4</td>
</tr>
<tr>
<td>long term depression</td>
<td>long term depression of brain cells</td>
<td>7.2x10^-4</td>
<td>0.075</td>
<td>5</td>
<td>ADCY5, CNR1, PRKCA, RYR3, ST8SIA4</td>
</tr>
<tr>
<td>long term depression</td>
<td>long term depression of neurons</td>
<td>0.0013</td>
<td>0.11</td>
<td>5</td>
<td>ADCY5, CNR1, DRD5, RYR3, ST8SIA4</td>
</tr>
<tr>
<td>assembly</td>
<td>assembly of olfactory cilia</td>
<td>0.0020</td>
<td>0.13</td>
<td>3</td>
<td>BBS10, PCDH15, VANGL2</td>
</tr>
<tr>
<td>long term depression</td>
<td>long term depression of corticostriatal neurons</td>
<td>0.0023</td>
<td>0.13</td>
<td>2</td>
<td>ADCY5, CNR1</td>
</tr>
<tr>
<td>positioning</td>
<td>positioning of cholinergic neurons</td>
<td>0.0023</td>
<td>0.13</td>
<td>2</td>
<td>SLIT1, SLIT2</td>
</tr>
<tr>
<td>contact repulsion</td>
<td>contact repulsion</td>
<td>0.0034</td>
<td>0.17</td>
<td>3</td>
<td>DCC, SEMA3A, SLIT2</td>
</tr>
<tr>
<td>chemotaxis</td>
<td>chemotaxis of neurons</td>
<td>0.0037</td>
<td>0.17</td>
<td>4</td>
<td>EPHB2, GFRA1, SEMA3A, SLIT2</td>
</tr>
<tr>
<td>development</td>
<td>development of optic chiasm</td>
<td>0.0068</td>
<td>0.22</td>
<td>2</td>
<td>SLIT1, SLIT2</td>
</tr>
<tr>
<td>size</td>
<td>size of lateral cerebral ventricle</td>
<td>0.0068</td>
<td>0.22</td>
<td>2</td>
<td>ANK2, NRG1 (includes EG:112400)</td>
</tr>
<tr>
<td>apoptosis</td>
<td>apoptosis of sensory neurons</td>
<td>0.0076</td>
<td>0.22</td>
<td>3</td>
<td>CDKN2D, HIPK2, KLF7</td>
</tr>
<tr>
<td>metabolism</td>
<td>fatty acid metabolism</td>
<td>0.010</td>
<td>0.22</td>
<td>5</td>
<td>DAB1, ERBB4, NRG1, SEMA3A, ST8SIA1</td>
</tr>
<tr>
<td>abnormal morphology</td>
<td>abnormal morphology of nervous system</td>
<td>0.013</td>
<td>0.22</td>
<td>2</td>
<td>CNR1, UHRF1</td>
</tr>
<tr>
<td>formation</td>
<td>formation of oligodendrocytes</td>
<td>0.013</td>
<td>0.22</td>
<td>2</td>
<td>ERBB4, NRG1</td>
</tr>
<tr>
<td>guidance</td>
<td>guidance of thalamocortical axons</td>
<td>0.013</td>
<td>0.22</td>
<td>2</td>
<td>SLIT1, SLIT2</td>
</tr>
<tr>
<td>migration</td>
<td>migration of GABAergic neurons</td>
<td>0.013</td>
<td>0.22</td>
<td>2</td>
<td>GFRA1, SLIT1</td>
</tr>
<tr>
<td>pathfinding</td>
<td>pathfinding of axons</td>
<td>0.013</td>
<td>0.22</td>
<td>2</td>
<td>DCC, EXT1</td>
</tr>
<tr>
<td>chemotaxis</td>
<td>chemotaxis of cells</td>
<td>0.014</td>
<td>0.22</td>
<td>5</td>
<td>EPHB2, FPFR2, GFRA1, SEMA3A, SLIT2</td>
</tr>
<tr>
<td>development</td>
<td>development of diencephalon</td>
<td>0.014</td>
<td>0.22</td>
<td>3</td>
<td>SIM2, SLIT1, SLIT2</td>
</tr>
<tr>
<td>long term depression</td>
<td>long term depression of synapse</td>
<td>0.018</td>
<td>0.22</td>
<td>3</td>
<td>CNR1, GRM7, MTOR</td>
</tr>
<tr>
<td>synthesis</td>
<td>synthesis of fatty acid</td>
<td>0.020</td>
<td>0.22</td>
<td>4</td>
<td>DAB1, ERBB4, NRG1, SEMA3A</td>
</tr>
<tr>
<td>synthesis</td>
<td>synthesis of lipid</td>
<td>0.020</td>
<td>0.22</td>
<td>6</td>
<td>CNR1, DAB1, ERBB4, NRG1, SEMA3A, ST8SIA1</td>
</tr>
</tbody>
</table>
Table 2. **Top 25 IPA Function Annotations for gC.** For each general function, the specific function annotation (FA) is shown together with its corresponding Fisher Exact Test (FET) P-value, the Benjamini-Hochberg (BH) multiple testing corrected P-value, the number of genes and the symbols of the genes responsible for the enrichment signal in the FA. The functions and FAs that remained significant in the IPA analysis of the genes that passed the more stringent criteria are underlined. The gene(s) in parentheses are those that emerged from the more conservative analysis.
### Table 3. Top 25 IPA Function Annotations for gF.

<table>
<thead>
<tr>
<th>Function</th>
<th>Function Annotation (FA)</th>
<th>FET P-value</th>
<th>BH P-value</th>
<th># Genes</th>
<th>Genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>synapatic fatigue</td>
<td>synapatic fatigue of synapse</td>
<td>0.0021</td>
<td>0.18</td>
<td>2</td>
<td>BDNF, SYN3</td>
</tr>
<tr>
<td>apoptosis</td>
<td>apoptosis of spinal cord cells</td>
<td>0.0041</td>
<td>0.18</td>
<td>4</td>
<td>BAX, BCL2, BDNF, FAM134B</td>
</tr>
<tr>
<td>inhibition</td>
<td>inhibition of neurons</td>
<td>0.0053</td>
<td>0.18</td>
<td>4</td>
<td>BDNF, NPPA, NPPB, NRG1</td>
</tr>
<tr>
<td>development</td>
<td>development of sensory nervous system</td>
<td>0.0062</td>
<td>0.18</td>
<td>2</td>
<td>BDNF, KIF1A</td>
</tr>
<tr>
<td>inhibition</td>
<td>inhibition of pyramidal neurons</td>
<td>0.0062</td>
<td>0.18</td>
<td>2</td>
<td>BDNF, NRG1</td>
</tr>
<tr>
<td>concentration</td>
<td>concentration of arachidonic acid</td>
<td>0.0062</td>
<td>0.18</td>
<td>2</td>
<td>DGKE, KNG1</td>
</tr>
<tr>
<td>endocytosis</td>
<td>endocytosis of synaptic vesicles</td>
<td>0.0067</td>
<td>0.18</td>
<td>4</td>
<td>AMPH, CABIN1, ITSN1, SNCA</td>
</tr>
<tr>
<td>damage</td>
<td>damage of cortical neurons</td>
<td>0.0067</td>
<td>0.18</td>
<td>3</td>
<td>BAX, BDNF, GRIN3A</td>
</tr>
<tr>
<td>quantity</td>
<td>quantity of central nervous system cells</td>
<td>0.0090</td>
<td>0.18</td>
<td>9</td>
<td>ACHE, AVPR1B, CCND2, EGFR, GDA, IGFBP6, LEF1, LHX5, TSC1</td>
</tr>
<tr>
<td>long-term potentiation</td>
<td>long-term potentiation</td>
<td>0.0092</td>
<td>0.18</td>
<td>13</td>
<td>B3GAT1, BDNF, CDH1, CHRNA7, CYP19A1, DGKE, LRP1, LRP8, NRG1, PLG, SNCA, VAV2, VAV3</td>
</tr>
<tr>
<td>apoptosis</td>
<td>apoptosis of dorsal root ganglion cells</td>
<td>0.0093</td>
<td>0.18</td>
<td>3</td>
<td>BAX, BCL2, FAM134B</td>
</tr>
<tr>
<td>mobilization</td>
<td>mobilization of Ca2+</td>
<td>0.0093</td>
<td>0.18</td>
<td>3</td>
<td>BDNF, PROK2, TRPV1</td>
</tr>
<tr>
<td>morphogenesis</td>
<td>morphogenesis of neurites</td>
<td>0.0097</td>
<td>0.18</td>
<td>22</td>
<td>ACHE, ATXN10, BDNF, BSN, CNTN4, CNTNAP2, CNTNA2, CTNND2, CYP19A1, EGFR, GDA, IGF1R, MAPK8, NRG1, PRKG1, PTPRM, RND1, SGK1, SLC18A3, SLIT1, SYNE1, UHMK1, (TLR7)</td>
</tr>
<tr>
<td>growth</td>
<td>growth of dendrites</td>
<td>0.0099</td>
<td>0.18</td>
<td>5</td>
<td>BDNF, CTNND2, CYP19A1, NRG1, SLIT1</td>
</tr>
<tr>
<td>morphology</td>
<td>morphology of dendrites</td>
<td>0.0099</td>
<td>0.18</td>
<td>5</td>
<td>BDNF, DISC1, GRIN2D, NPA53, NRG1</td>
</tr>
<tr>
<td>quantity</td>
<td>quantity of acetylcholine</td>
<td>0.012</td>
<td>0.18</td>
<td>2</td>
<td>ACHE, SLC18A3</td>
</tr>
<tr>
<td>survival</td>
<td>survival of dorsal root ganglion</td>
<td>0.012</td>
<td>0.18</td>
<td>2</td>
<td>BAX, BDNF</td>
</tr>
<tr>
<td>synaptogenesis</td>
<td>synaptogenesis of brain cells</td>
<td>0.012</td>
<td>0.18</td>
<td>2</td>
<td>BDNF, CYP19A1</td>
</tr>
<tr>
<td>long-term potentiation</td>
<td>long-term potentiation of granule cells</td>
<td>0.012</td>
<td>0.18</td>
<td>2</td>
<td>BDNF, DGKE</td>
</tr>
<tr>
<td>sensitization</td>
<td>sensitization of neurons</td>
<td>0.012</td>
<td>0.18</td>
<td>2</td>
<td>BDNF, KNG1</td>
</tr>
<tr>
<td>migration</td>
<td>migration of GABAergic neurons</td>
<td>0.012</td>
<td>0.18</td>
<td>2</td>
<td>GFRA1, SLIT1</td>
</tr>
<tr>
<td>quantity</td>
<td>quantity of astrocytes</td>
<td>0.012</td>
<td>0.18</td>
<td>4</td>
<td>ACHE, EGFR, IGFBP6, TSC1</td>
</tr>
<tr>
<td>cell viability</td>
<td>cell viability of motor neurons</td>
<td>0.015</td>
<td>0.18</td>
<td>5</td>
<td>BAX, BCL2, BDNF, GFRA1, REG3G</td>
</tr>
<tr>
<td>loss</td>
<td>loss of motor neurons</td>
<td>0.016</td>
<td>0.18</td>
<td>3</td>
<td>BCL2, BDNF, GFRA1</td>
</tr>
<tr>
<td>quantity</td>
<td>quantity of nerve ending</td>
<td>0.016</td>
<td>0.18</td>
<td>3</td>
<td>BDNF, SLC18A3, SNCA</td>
</tr>
</tbody>
</table>
Table 3. **Top 25 IPA Function Annotations for gF.** For each general function, the specific function annotation (FA) is shown together with the corresponding Fisher Exact Test (FET) P-value, the Benjamini-Hochberg (BH) multiple testing corrected P-value, the number of genes and the symbols of the genes responsible for the enrichment signal in the function annotation. FAs that remained significant after excluding BDNF are in bold. The functions and FAs that remained significant in the IPA analysis of the genes that passed the more stringent criteria are underlined. The gene(s) in parentheses are those that emerged from the more conservative analysis.