Interpreting a migraine GWAS using gene expression in healthy human brain



PRDM16

othelial cells roglia

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Summary

Migraine is a common brain disorder, with a heritability of 50%. Genome-wide association studies have identified several loci, but interpretation remains challenging. We integrated migraine GWAS data with spatial gene expression data of adult brains from the Allen Human Brain Atlas, to identify

Migraine GWAS

GWAS data¹ from 23,285 migraine patients - 95,425 controls.

Calculate p-values per gene with GATES²: - LD and gene size corrected. - SNPs within 15 kb flanks.

Healthy brain gene expression

Gene expression data³ from the Allen Human Brain Atlas.

Expression in 3702 samples from six healthy human donors, covering most

specific brain regions and molecular pathways involved in migraine.

We used two complementary methods. First, we clustered all genes into co-expression modules and identified those associated with migraine. Second, we constructed local co-expression networks around high-confidence migraine genes. Both approaches converge on functions and anatomy.

Define migraine genes: p-value per gene with GATES - "High confidence genes"

- Bonferroni corrected p < 0.05
- "Candidate genes" uncorrected p < 0.05

of the brain.

Use these samples to calculate spatial co-expression between genes.

If genes are co-expressed, they share a spatial expression pattern.

TRPM8

Approach 1

Using co-expression data from the healthy brains, we clustered all genes into 18 co-expression modules.

All modules were tested for enrichment in migraine "candidate genes". Five modules of interest were identified (A - E).

These modules are involved in:

- Neurotransmission, protein catabolism and mitochondria in the cortex.
- Transcription regulation in the cortex and

Approach 2

biological functions.

We selected the 14 "high confidence migraine" genes" to serve as seeds in local co-expression networks.

Each of these genes was connected to its most co-expressed genes in the healthy brains.

The network shows considerable overlap with modules A, B and D of Approach 1. It also points to the same anatomical regions and

cerebellum.

 Oligodendrocytes and mitochondria in subcortical areas.





Module

Module

Ω

Module



1,556 genes / 205 candidate genes $p = 9.22 \times 10-4$ LD corrected $p = 5.47 \times 10^{-4}$

Function EASE Integration of energy metabolism 3.84 Mod.-dep. protein catabolic process 2.61 2.57 Proteasome 2.09 Synapse Voltage-gated cation channel activity 2.08

1,595 genes / 198 candidate genes p = 0.015 LD corrected $p = 7.18 \times 10^{-3}$

unction	EASE
luclear lumen	16.46
inc finger transcription factor	13.85
ranscription	13.19
inc ion binding	9.18
Chromatin modification	8.02

497 genes / 67 candidate genes p = 0.020LD corrected $p = 7.77 \times 10^{-3}$

p = 0.024

Function

Apoptosis

Mitochondrial part

Mitochondrion

EASE Function 3.21 Membrane fraction Mod.-dep. protein catabolic process 3.13 Purine ribonucleotide binding 2.61



FHL5

Cerebellar cortex formation 1.71 Regulation of synaptic plasticity

1.70







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