Functional gene networks reveal distinct mechanisms segregating in migraine families

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Migraine is the most common neurological disorder worldwide and it has been shown to have complex polygenic origins with a heritability of estimated 40–70%. Both common and rare genetic variants are believed to underlie the pathophysiology of the prevalent types of migraine, migraine with typical aura and migraine without aura. However, only common variants have been identified so far. Here we identify for the first time a gene module with rare mutations through a systems genetics approach integrating RNA sequencing data from brain and vascular tissues likely to be involved in migraine pathology in combination with whole genome sequencing of 117 migraine families. We found a gene module in the visual cortex, based on single nuclei RNA sequencing data, that had increased rare mutations in the migraine families and replicated this in a second independent cohort of 1930 patients. This module was mainly expressed by interneurons, pyramidal CA1, and pyramidal SS cells, and pathway analysis showed association with hormonal signalling (thyrotropin-releasing hormone receptor and oxytocin receptor signalling pathways), Alzheimer’s disease pathway, serotonin receptor pathway and general heterotrimeric G-protein signalling pathways. Our results demonstrate that rare functional gene variants are strongly implicated in the pathophysiology of migraine. Furthermore, we anticipate that the results can be used to explain the critical mechanisms behind migraine and potentially improving the treatment regime for migraine patients.

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Abbreviations: MAF = minor allele frequency; MTA = migraine with typical aura; WGCNA = weighted gene co-expression network analysis; WGS = whole genome sequencing

Introduction

Migraine is a common neurovascular disorder that is characterized by a unilateral throbbing head pain, accompanied by sensitivity to light and sound and/or nausea/vomiting. It is often accompanied by neurological disturbances (the aura) (Headache Classification Committee of the International Headache Society (IHS), 2018; Szabó et al., 2018). The prevalent types of migraine, migraine with typical aura (MTA) and migraine without aura are considered complex polygenic disorders with a heritability between 40% and 70% (Sutherland and Griffiths, 2017). Scores of common genetic variants have been associated with migraine explaining a part of the genetic variance (Gormley et al., 2016). It has also been shown that the polygenic risk score is higher...
in families with MTA and migraine without aura; however, it is believed that rare causal variants could also contribute to the genetic variance of migraine (Hansen et al., 2017; Gormley et al., 2018). Such rare causal variants can be detected in families with mendelian-like segregating pattern of migraine, using e.g. whole genome sequencing (WGS). It is encouraging that family studies have identified multiple rare causal variants for a rare dominant form of migraine, familial hemiplegic migraine (Ophoff et al., 1996; Dichgans et al., 2005; Marini et al., 2012; Barros et al., 2013). However, as the prevalent types of migraine are polygenic and do not follow a complete dominant inheritance pattern, we hypothesize that genes with rare variants of small or medium effect size are aggregating in mechanisms specific for migraine. Such mechanisms can be identified in gene networks identified, for example, by RNA sequencing of migraine-relevant tissues in combination with WGS.

Weighted gene co-expression network analysis (WGCNA) (Zhao et al., 2010) has been useful in identifying such gene networks, which has the potential of pointing towards biologically and clinically relevant disease mechanisms (Eising et al., 2017; Feng and Wang, 2017; Liao et al., 2017; Wang et al., 2017). Gene expression is highly tissue-specific, thus co-expression networks of tissues likely to be involved in migraine could suggest migraine-specific networks. The pathophysiology of migraine is not completely understood but it is widely accepted and shown in several studies that the trigeminal ganglion, the visual cortex and the vasculature are involved in critical mechanisms of migraine (Zhang et al., 2010, 2011; Coppola et al., 2013; Brighina et al., 2015; Gormley et al., 2016; Shelukhina et al., 2017; Gaist et al., 2018). The trigeminal ganglion is a cluster of nerve cells controlling sensory input from the face and, for example, due to release of calcitonin gene-related peptide (CGRP) and expression of serotonin 5-HT3α, a treatment target of migraine (Potrebic et al., 2003; Messlinger and Russo, 2018). The visual cortex is involved in processing of visual information, and cortical spreading depression in visual cortex explains the aura (Marzoli and Criscuoli, 2017; Charles, 2018). The cardiovascular system has, in addition to specific brain tissues, been linked to migraine (Pickrell et al., 2016). This was recently stipulated as aorta, as common genetic variants associated with migraine were significantly higher expressed in aorta (Gormley et al., 2016).

In this study, we used a systems genetics approach integrating rare genetic variants segregating in migraine families with gene expression networks. We first identified gene networks using WGCNA applied to RNA sequencing data from human aorta (tissue), trigeminal ganglion (tissue) and visual cortex (single nuclei). Second, we integrated rare mutations from our WGS data with gene networks to pinpoint networks having rare variants segregating with migraine, suggesting distinct involvement in migraine mechanisms. Third, these results were replicated in an independent cohort of migraineurs without familial incidences of migraine. The complementary results revealed—for the first time—the critical role of rare gene variants in migraine heritability.

Materials and methods

The study design focused on aorta, trigeminal ganglion and visual cortex as they are key tissues in the pathophysiology of migraine (Zhang et al., 2010, 2011; Coppola et al., 2013; Brighina et al., 2015; Gormley et al., 2016; Shelukhina et al., 2017; Gaist et al., 2018), aiming to identify the pathways involved in the migraine mechanisms (Fig. 1). We constructed co-expression networks of aorta, trigeminal ganglion and visual cortex using RNA sequencing data, for identification of tissue-specific modules using WGCNA. We then integrated data on rare functional variants segregating with migraine, derived from WGS data of 874 individuals from 117 families with migraine, to extracted modules associated with migraine. We aimed to replicate the modules in patients with sporadic migraine (n = 1930) and population control subjects (n = 2500). We then performed functional annotation to map the pathways/networks and cell types involved in migraine.

Subjects

We recruited 117 families with Mendelian-like segregation of migraine. These families consisted of 874 subjects, 262 of which were diagnosed with MTA, 213 with migraine without aura, 145 with both MTA and migraine without aura, and 234 familial members with no history of migraine. Initially, a migraine patient, i.e. proband, was recruited from the Danish Headache Center and, subsequently, family members were recruited, enabling segregation analysis. The replication cohort consisted of 1930 sporadic migraine patients, 312 of whom were diagnosed with MTA, 1087 were diagnosed with migraine without aura, and 531 were diagnosed with both MTA and migraine without aura; all were recruited from the Danish Headache Center. All participants were assessed by a trained physician or a senior medical student trained in headache diagnostics using a semi-structured interview. The interview was designed by the previous head of classification committee Prof. Jes Olesen to enable diagnosis according to the International Classification of Headache Disorders second edition (ICHD-3) (Olesen, 2006). The study was approved by the Ethics Committee (H-2-2010-122), reported and approved by the data protection agency, and written informed consent was obtained from all participants.

For population controls, we used sequencing data from the 1000 Genomes project (1000G) (Auton et al., 2015). We used the 1000G phase 3 data (VCF files), which included 2500 individuals who were both WGS and whole exome sequenced (WES). When restricting the 1000G cohort to non-Finish Europeans (NFE), 403 individuals were analysed. The NFE classification was provided by 1000G.

Whole genome sequencing

Preparation of samples

Genomic DNA extraction was prepared from whole blood using Illumina TruSeq DNA kit according to the manufacturers’ instructions. In short, 1 μg genomic DNA was isolated from frozen blood and fragmented into 300–400 bp using a Covaris
E220 instrument, followed by end repair. AMPure XP magnetic purification beads were then used for size selection followed by PCR enrichment (10 cycles) using appropriate primers. Subsequently, 3'-adenylation and ligation of sequencing adaptors with a T nucleotide overhang was performed and again followed by AMPure magnetic purification. The sequencing libraries were then analysed with regards to quality and concentration using the LabChip® GX (96 samples) instrument from Perkin Elmer. Sequencing libraries were diluted and stored at –20°C. The optimal cluster densities, library insert size, duplication rates and library diversities were assessed by multiplexing on an Illumina MiSeq instrument. All steps in the workflow were monitored using an in-house laboratory information management system (LIMS) with barcode tracking of all samples and reagents.

Whole genome sequencing

Sequencing libraries were hybridized to the surface of paired-end flow cells using the Illumina cBot. Paired-end sequencing-by-synthesis was performed on an Illumina HiSeq X instrument. Read lengths used were 2 × 150 cycles of incorporation and imaging. Real-time analysis involved conversion of image data to base-calling in real-time.

Reference genome and alignment

The human genome assembly GRCh38 was used as reference sequences map reads. The reference did not include alternate assemblies (GCA_000001405.15_GRCh38_no_alt_analysis_set.fna) in addition to sequences determined to represent common contaminants in the sequencing pipeline. Common contaminants in the sequencing pipeline are the sequences of the bacteriophage PhiX (Enterobacteria_phage_phiX174_sensu_lato_uid14015/NC_001422.fna), the bacteria Ralstonia Pickettii (Ralstonia_pickettii_12D_uid58859) and two sequences from the human microbiome (Coprobacillus_D7_uid32495/NZ_EQ999972.fna and Coprobacillus_D7_uid32495/NZ_EQ999922.fna).

Raw sequences were aligned against the reference with BWA v.0.7.10 (Li and Durbin, 2009). The sequences in the BAM files were realigned around indels with GenomeAnalysisTK-Lite/ 2.3.9 (McKenna et al., 2010) using a public set of known indels. We marked PCR duplicates with Picard tools v.1.117.

Quality control and variant calling

After alignment to reference genome, BAM files were analysed and removed from the analysis based on following criteria: mean base quality >25, per cent marked duplicate >50, mean n per read >30, per cent mapping quality below 20 >11, per cent reads unmapped >40, per cent both reads in a pair unmapped >40, per cent first read in a pair unmapped >40, per cent second read in a pair unmapped >40. Following filtering, variants were called, which resulted in 5 017 453 ± 2607 variants per individual and 697 044 ± 3244 variants with minor allele
frequency (MAF) < 5% (Supplementary Fig. 1B and C) and a mean sequencing depth of 32.94 ± 0.22 (Supplementary Fig. 1A).

WGS and variant calling were performed in collaboration with deCODE genetics®.

**RNA sequencing data**

RNA sequencing data of aorta (n = 247 individuals) were downloaded from GTEx using the R-package recount (Collado-Torres et al., 2017). Sequencing and alignment are described at http://gtexportal.org. In short, RNA sequencing was performed using the Illumina TruSeq library construction protocol, and sequenced on the Illumina Hiseq2000 or HiSeq2500, generating 76-bp paired-end reads. Reads were aligned to the human reference genome (GRCh38) using STAR v.2.5.3a based on the GENCODE v.26 annotation. Read counts and TPM values were produced with RNA-SeQC.

RNA sequencing data of human trigeminal ganglia (n = 16 individuals) was downloaded from NCBI’s Sequence Read Archive (SRX3020382). Messenger RNA was isolated from small subregions of 16 human trigeminal ganglia, as described previously (LaPaglia et al., 2018). In short, non-stranded cDNA libraries were prepared using the Illumina TruSeq RNA Library Preparation Kit v2, and then sequenced on the Illumina HiSeq2500, generating 125-bp paired-end reads. The quality of the fastq-files was checked using the NGSQC toolkit and low quality reads were filtered out according to default settings (cut-off read length of 70% and a cut-off quality score of 20). On average, per sample, 85.2% of the reads were of high quality resulting in on average 26.8 million reads [standard deviation (SD) = 1.9 million reads]. Filtered fastq files were aligned to the human reference genome (Release 32, GRCh38.p13) using the pseudo-aligner Kallisto v.0.46.0. Length-scaled transcripts per million were imported into R using tximport v.3.10.

RNA sequencing data of the human primary visual cortex were downloaded from Allen Human Brain Map (n = 8998) and technical details are described in the technical white paper: transcriptomics (Allen Brain Institute, 2018). In short, 8998 single nuclei were isolated from six cortical layers of visual cortex (n = 8 individuals). The Nextera XT kit (Illumina) was used to prepare cDNA libraries. Single nucleus libraries were then sequenced on the Illumina HiSeq2500, generating 50-bp paired-end reads. Reads were aligned to the human reference genome (GRCh38.p2) using STAR v.2.5.3. Final results included quantification of exon and intron counts at gene level. We merged exon and intron data by summing exon and intron counts at gene level.

For all tissues, we included only protein-coded genes, using a gene list extracted from BioMart (GRCh38.p18). This resulted in counts of 19747, 18299 and 16665 genes in trigeminal ganglion, visual cortex and aorta, respectively.

**Construction of co-expression networks**

A co-expression network was constructed of each tissue, i.e. aorta, trigeminal ganglion and visual cortex. Genes for network construction were selected based on their connectivity (sum of correlations of a gene with all other genes); the 5000 most highly connected genes per tissue were retained. A network was constructed based on the default pipeline of WGCNA (Zhang et al., 2005; Langfelder and Horvath, 2008). To ensure scale-free topology, the correlation network was raised to a power (beta) to reach a scale-free topology measure (R2) > 0.8. To ensure a stable and highly reliable network, the adjacency matrix and consequently dissimilarity topological overlap matrix (TOM) were calculated 100 times by leaving 10% of the samples out every time. The average dissimilarity TOM was calculated from the 100 networks. Next, a gene dendrogram was created and modules containing at least 30 genes were detected using the dynamical tree cutting algorithm. Modules are defined as clusters of highly co-expressed genes. Modules were validated using a permutation procedure as described by Iancu et al. (2012). For each network, we randomly selected groups of genes with the same size of the detected modules in the network and computed the average topological overlap. All modules in the aorta, trigeminal ganglion and visual cortex networks had a significantly higher topological overlap than the random modules, except for the ‘grey’ module (which includes unassigned genes). This confirmed that we detected modules with statistical and potentially functional relevance for the tissues.

**Detection of migraine-associated modules**

**Detection of mutated genes in migraine families**

We used the pedigree variant annotation, analysis and search tool (pVAAST) v.2.2.0 (Hu et al., 2014) to detect genes with rare functional mutations segregating with migraine. We performed a test for each gene in the modules for each of the 117 families.

Genes were identified with pVAAST based on the following criteria: (i) genes were identified based on the penetrance level (0.9–1) in each family; and (ii) genes were identified based on the functionality of variants only giving rise to frameshift deletions, frameshift insertions, non-framen shift deletions, non-frame shift insertions, non-synonymous single nucleotide polymorphisms (SNPs), splice site mutations, stop-gain and stop-loss mutations. We refer to these functional variants as mutations from here on. Third, genes were identified based on mutations having a MAF < 5% in the genome aggregation database (gnomAD) v.2. See Supplementary Fig. 3A–C for MAF distribution of all variants in the three tissues analysed. The rationale for choosing a MAF < 0.05% was based on the prevalence of migraine being 1:5 to 1:6, biallelic, a genetic heterogeneity of 1, several scenarios of allelic heterogeneity (between 0.01 and 0.05) gave rise to MAF up to 5%, thus a MAF cut-off of 5% was chosen. The cut-off was calculated using the allele frequency app https://www.cardiodb.org/allelefrequency (Whiffin et al., 2017).

Genes with rare functional mutations are hereafter termed mutated genes. Genes were termed non-mutated when at least one variant could be detected that was not a rare functional variant.

**Genic intolerance analysis of modules**

The mutation tolerance is gene dependent, thus we calculated the mean residual variation intolerance score (RVIS) (Petrovski et al., 2013) for genes with a rare functional mutation (mutated genes) and for genes without rare functional mutations (non-mutated genes) per module. A one-sided t-test was performed to
exclude modules where the mutated genes had a significantly higher mean RVIS than the non-mutated genes \((P < 0.05)\).

**Detection of migraine-associated modules in families**

For each tissue, modules with more mutated genes than expected by chance were detected by a one-tailed Fisher’s exact test, i.e. the number of mutated genes versus non-mutated genes in a module against number of mutated genes versus non-mutated genes in all other modules. We only included genes in the Fisher’s exact test in which genetic variations were detectable. Therefore, the number of genes included in Fisher’s exact test for familial migraine were: 4894 in aorta, 4652 in trigeminal ganglion, and 4879 in visual cortex. \(P\)-values were adjusted using Bonferroni correction for the number of modules (trigeminal ganglion: \(n = 39\), visual cortex: \(n = 8\), aorta: \(n = 16\)) and defined as migraine-associated when \(P_{\text{adj}} < 0.05\).

**Replication of migraine-associated modules in sporadic migraine**

Using an independent cohort of 1930 sporadic migraine patients, i.e. with no known history of first-degree relatives with migraine, and using the 1000G control cohort, we assessed the migraine-associated modules found in the migraine family cohort using ANNOVAR (v.2018apr16). ANNOVAR was run using VCF files to annotate genetic variants in mRNAs and to annotate MAF of those genetic variants. We used refSeq and gnomAD v.2 as input databases for ANNOVAR. Module genes with a MAF < 5% were analysed for mutations as defined above. See Supplementary Fig. 3D–F for MAF distribution of mutations. Modules with more mutated genes than expected by chance were determined using one-tailed Fisher’s exact test, i.e. the number of mutated genes versus non-mutated genes in a module against number of mutated genes versus non-mutated genes in all other modules. We only included genes in the Fisher’s exact test in which we could detect genetic variation. Therefore, the number of genes included in the test were for sporadic migraine: 4871 in trigeminal ganglion, 4973 in visual cortex and 4956 in aorta. For 1000G the gene counts were: 3201 in trigeminal ganglion, 2937 in visual cortex and 3251 in aorta. To avoid biases derived from different data processing and batches, the Fisher’s tests were carried out separately for cases and controls. Modules that had significantly more mutated genes in cases but not controls were considered replication.

**Functional annotation of migraine-associated modules**

**Visualization**

The Protein-Protein-interaction network was depicted using stringApp v.1.5.0 in Cytoscape v.3.7.2 (Shannon et al., 2003) assessing the protein products and interaction thereof having the threshold for the interaction confidence score > 0.4. Protein interaction data v.11 from www.string-db.org were used as input for Cytoscape.

**Pathway analysis**

We carried out pathway analysis using PANTHER over-representation test (data version released 20190711) (Mi et al., 2013). The total set of genes used for network construction of the tissue of interest was used as reference. Pathways were tested for over-representation using the binomial test and \(P\)-values were adjusted for multiple-testing using FDR correction and defined as significant in case \(P_{\text{adj}} < 0.05\).

**Enrichment analysis in single cell RNA sequencing data**

We used two different approaches to investigate modules with increased mutation burden segregating in families. First, we used single cell RNA sequencing data from mouse cortex and hippocampus for cell-type enrichment testing (Zeisel et al., 2015), allowing for differentiation between six different cell types: astrocytes, interneurons, endothelial cells, microglia, oligodendrocytes, pyramidal CA1 neurons and pyramidal SS neurons. Enrichment analysis was carried out using the Expression Weighted Celltype Enrichment (EWCE) R package (Skene and Grant, 2016) with 10000 bootstraps for genes in the significant migraine-associated modules.

**Data availability**

WGS data are considered personal and sensitive data by the scientific ethical committee, thus data can only be shared as part of material agreed upon reasonable request.

**Results**

We constructed gene co-expression networks for three tissues known to be involved migraine aetiology: the aorta, the trigeminal ganglia, and the visual cortex. For each tissue we selected the 5000 most highly connected genes for network construction; 961 genes were present in all three networks (Fig. 2A and Supplementary Table 1). From the derived co-expression networks, we identified 34 modules in the aorta network, 64 modules in trigeminal ganglia and eight modules in visual cortex (Supplementary Fig. 2 and Supplementary Table 1). Though the aorta and trigeminal ganglia networks are based on tissues, the visual cortex network is based on single nuclei data and, therefore, modules may represent subgroups of cell types. Cell type enrichment analysis showed that, for example, the green module was over-represented by genes expressed in interneurons, though other modules (e.g. blue) were covered by different cell types (Fig. 2E).

We used WGS data from our cohort of families with Mendelian-like segregation pattern of migraine, to identify migraine-associated modules, with over-representation of mutated genes. We excluded gene modules that had over-representation of highly mutation-tolerant genes, i.e. type I error (Supplementary Fig. 4). From the aorta network, 16 modules remained for analysis, 39 in trigeminal ganglia and eight in visual cortex.

We identified 313 mutated genes in aorta, 875 mutated genes in trigeminal ganglia, and 1243 mutated genes in visual cortex, of which 13 were present in all three networks (Fig. 2A and B). We found no significant overlap of mutated genes between the three networks (Fig. 2C). The rare mutations consisted primarily of non-synonymous SNPs.
while 0.54–5.43% are high impact mutations (frameshift mutations, splice site mutations and stop-loss/stop-gain mutations) (Fig. 2D).

Selection of migraine-associated modules

As hypothesized, genes with rare mutations are more likely to aggregate in pathways specific for migraine mechanisms. Such genes are likely to be enriched within our constructed networks, based on the expression profile of migraine-relevant tissue. In the aorta network we found one migraine-associated module (Fig. 3A, darkgreen), in the trigeminal ganglion network we found five migraine-associated modules (yellow, salmon, honeydew1, darkturquoise and darkslate-blue), and in the visual cortex network we found three migraine-associated modules (blue, brown, yellow) (Fig. 3A).

We were able to replicate the brown module (visual cortex), showing an increased burden of mutated genes in our sporadic migraine cohort and not in the 1000G cohort. We were not able to replicate any of the other modules, although the yellow module (trigeminal ganglia) was significant in the sporadic migraine it was also significantly increased in the 1000G cohort (Fig. 3B). Restricting the 1000G cohort to non-Finish Europeans did not affect the results (Supplementary Fig. 5A–C).

We analysed the visual cortex brown module and the trigeminal ganglion yellow module on a protein level.

(86.59–90.13%), while 0.54–5.43% are high impact mutations (frameshift mutations, splice site mutations and stop-loss/stop-gain mutations) (Fig. 2D).
The majority (82.5%) of the proteins in the visual cortex brown module were participating in the network, suggesting a strong functional module network (Fig. 4A). The trigeminal ganglion yellow module was divided into three smaller networks covering 5.7–32.1% of the proteins, indicating a less strong functional module network (Fig. 4B).

**Functional annotation of migraine-associated modules**

Cell type enrichment analysis of the migraine-associated module (visual cortex brown) showed that the module was mainly expressed by interneurons, pyramidal CA1, and pyramidal SS cells (Fig. 2E). Furthermore, five pathways were over-represented: the thyrotropin-releasing hormone receptor (TSHR) signalling pathway, oxytocin receptor (OXTR) mediated signalling pathway, Alzheimer’s disease amyloid secretase pathway, serotonin receptor (5HT2R) signalling pathway, and heterotrimeric G-protein signalling pathway-Gq alpha and Go alpha mediated pathway (Table 1). The distribution of mutated genes co-segregating with migraine across families for the module showed that 71 genes have mutations in one family whereas 126 genes harbour functional mutations in two or more families (Fig. 5). Several genes were mutated frequently among families. The top three genes included ATXN1, mutated in 28 families; FAM153B, mutated in 19 families; and CACNA1B, mutated in 16 families.
Discussion

We present a systems genetics approach integrating WGS data from 117 families with Mendelian-like segregation patterns of migraine with RNA sequencing data from tissues involved in migraine aetiopathology: trigeminal ganglia, visual cortex, and aorta. After replication in an independent cohort of 1930 sporadic cases of migraine, we identified one module with a significant accumulation of mutated genes. The results propose distinct mechanisms affected in migraine, including the TSHR signalling pathway, OXTR signalling pathway, Alzheimer’s disease amyloid secretase pathway, 5HT2 type receptor mediated signalling pathway and Heterotrimeric G-protein signalling pathway-Gq alpha and Go alpha mediated pathway. Further, we were able to suggest specific genes with mutations to be involved in migraine aetiology.

The co-expression approach has been applied to migraine previously. Focusing on common risk variants and using different brain tissues (Eising et al., 2016). The authors allocated migraine-associated genes to modules across brain tissues, showing that oligodendrocytes and gene transcription regulation may play a causal role in migraine. We have shown a similar approach with two important differences. First, our networks are constructed using human RNA sequencing data from the tissues probably involved in migraine aetiopathology, trigeminal ganglia, visual cortex, and aorta. Second, we integrated the co-expression networks with mutations found segregating in a large family cohort.

WGCNA is a well-established method for studying biological networks and diseases. We used it to identify gene modules in three migraine-relevant tissues. Like WGCNA, WGS is also a well-known resource in disease research. Over recent years, the cost of having a genome sequenced has dropped substantially such that it is now possible to sequence entire cohorts (Hayden, 2014). WGS has an advantage over, for example, genotyping since rare variants and de novo variants are included in the sequencing. We have—for the first time—utilized both methods in an attempt to reveal more of the defective biological mechanisms leading to migraine. Common variants do not explain 100% of the heritability of migraine (Gormley et al., 2016). Integrating RNA sequencing data via WGCNA with WGS, we have closed some of the gap between how much of the phenotype can be explained by common variants and now by rare mutations (MAF < 0.05%).

Serotonin and the serotonin signalling pathway have, for many years, been known for their implications in migraine. Serotonin levels are believed to increase during migraine attacks (Sicuteri et al., 1961) and the family of serotonin receptor agonists—triptans—are the preferred migraine treatment (Dussor, 2014). Serotonin binds to serotonin receptors, which comprise both metabotropic and ionotropic receptors. The two families of metabotropic serotonin receptors: 5HT1 and 5HT2, have both been associated with migraine (Hill and Reynolds, 2007; Ferrari et al., 2010; Nelson et al., 2010; Thompson et al., 2012; Yücel et al., 2016) as well as the serotonin transporter (Ogilvie et al., 1998; Marziniak et al., 2005; Bayerer et al., 2010). We directly associate 5HT2 with migraine in the visual cortex brown module as well as the G-protein-mediated signalling pathway including G, which signals downstream of the 5HT1 receptor. Moreover, we found an association of the visual cortex brown module gene-set to neurons in our EWCE analysis. Cortical neuronal activity initiates and propagates cortical spreading depression (Leao, 1944), which has been linked to serotonergic activity in functional studies (Supornsilpchai et al., 2006; Guedes et al., 2017). We suggest that the migraine phenotype may be caused by mutated genes in the serotonin signalling pathway as well as the downstream G-protein coupled signalling pathway.

In addition to serotonergic neurons, glutamatergic neurons and synapses have also been linked to migraine through different association studies where SNPs in genes related to glutamate transport or signalling have been detected.
Glutamate levels have been increased in migraineurs (Ferrari et al., 1990; Martinez et al., 1993; Campos et al., 2013). Glutamate plays a central role in cortical spreading depression as an activator and propagator and inversely, inhibition of glutamate receptors impairs initiation and propagation of cortical spreading depression (Lauritzen and Hansen, 1992; Peeters et al., 2007; Zhou et al., 2013). In our analysis, we detected three genes (ATXN1, FAM153B and CACNA1B) that were mutated in >15 families. CACNA1B and ATXN1 are both involved in glutamate signalling. CACNA1B encodes a subunit in a voltage-gated calcium channel, which is expressed at the presynapse and involved in release of glutamate from the presynapse (Bunda et al., 2019). ATXN1 encodes a DNA binding protein, which is able to regulate expression of the metabotropic glutamate receptor 1 (mGluR1) at the postsynaptic membrane (Notartomaso et al., 2013). Thereby the glutamatergic signalling is altered on both sides of the synaptic cleft in our migraine patients and may contribute to the migraine pathophysiology. CACNA1B is a member of a family of genes, which includes CACNA1A, that has been linked to a monogenic subtype of MTA, namely familial hemiplegic migraine (FHM) (Ophoff et al., 1996). FHM is a more severe phenotype than MTA and patients with FHM can experience ataxia symptoms. Ataxia is a complex disease and multiple genes have been linked to the disorder including ATXN1. The knowledge regarding FAM153B is limited and no function of the encoded protein has been reported. According to the GTEX data, FAM153B is highly expressed in the brain and likely has a vital function in neurons, which may be affected by mutations (Lonsdale et al., 2013).

Oxytocin and thyrotropin are both hormones that regulate the activity of OXTR and TSHR. Both receptors are G-protein coupled receptors expressed throughout the brain; however, TSHR is expressed in lower amounts than OXTR in the brain according to GTEX data (Lonsdale et al., 2013; Vafaviinge et al., 2020). Oxytocin has been the subject of migraine research as it is a strong analgesic in neuroinflammation (Yu et al., 2003; Yang et al., 2007; Engle et al., 2012; Kubo et al., 2017). Moreover, oxytocin is increased during pregnancy, which inversely correlates to the decrease in headaches and migraine of pregnant females (Callaghan, 1968; Kuwabara et al., 1987; Goadsby et al., 2008; Hoshiyama et al., 2012). Oxytocin levels also increase during intercourse and orgasms for both males and females. For the latter, it is reported that 47% experience relief from migraine during sex (Carmichael et al., 1987; Evans and Couch, 2001). Clinical testing of oxytocin in migraine patients led to a long term (24 h) effect on pain after dosing. In the course of 8 weeks, oxytocin had an effect on the migraine frequency (Tzabazis et al., 2017). Thyrotropin is less studied with respect to migraine compared to oxytocin. However, high levels of thyrotropin were associated with lower occurrences of headache (Hagen et al., 2001) and low levels of thyrotropin may be associated with a prolonged and refractory clinical course of migraine (Starikova et al., 2019). Taken together, hormone receptor misregulation seems to play a role in migraine. Our data point to a dysregulation in the hormone receptor pathways, which most likely derives from mutations affecting proteins active downstream from oxytocin and thyrotropin receptors.

Finally, we detected the Alzheimer’s disease amyloid secre- tase pathway in the visual cortex brown module. It has been suggested that migraine may be associated with mild types of cognitive changes (Araújo et al., 2012) but there is no clear evidence showing whether or not migraine is a risk factor for Alzheimer’s disease (Breiter et al., 1991; Wang et al., 2018; Lee et al., 2019; Morton et al., 2019).

**Strengths and limitations**

In this study we used a novel approach by combining gene-gene interaction networks based on RNA sequencing data with rare mutations found in families using WGS data. The co-expression networks enable investigation of biological mechanisms in migraine-relevant tissues. The construction of those networks is affected by the origin and quality of the RNA sequencing data, i.e. inter-tissue comparison is not possible because of, for example, their origin, bulk versus single cell, and sequencing depth. The modules revealed in the aorta and trigeminal ganglia data are driven by differences in gene expression levels between individuals, while in the visual cortex network, using single nuclei data, modules are driven by differences in cell types. Interpretation of pathways detected in visual cortex, therefore, needs to be done with caution. Similarly, thresholds applied to mutation calling affect results. We have analysed mutations with a MAF < 0.05 in both familial and sporadic migraine. The recent migraine GWAS included variants with a MAF as low as 0.01 (Gormley et al., 2016). Our study may, therefore, overlap with respect to these variants. However, we believe it is a strength to include variants with a MAF up to 0.05 because of the high prevalence of migraine in the population.

The differences in the number of genes having genetic variants in the families was marginal, which is also reflected by the mean coverage of the genome. The difference between the replication cohorts most likely derives from different sample handling and sequencing, hence the separate assessment of the sporadic migraine cohort and 1000G cohort. Thus, a more ideal comparison of having controls from the same population and process together would have been preferred.

Our familial migraine cohort has a large fraction of migraine patients with aura. This is due to the initial recruiting of our patients and may be considered as a limitation. However, our replication cohort of sporadic migraine has a small fraction of MTA patients compared to migraine without aura patients. Despite the fact that there are no differences in clinical presentation between familial migraine and sporadic migraine, our results may be slightly biased towards MTA since we initially analysed our familial cohort (Ravn et al., 2019). If more familial migraine without aura patients were included in future studies resembling our
study, it could perhaps result in more gene modules implicated in the migraine biology.

Conclusion
Migraine-associated mechanisms derived from a visual cortex module exist and are likely to contribute to the migraine pathophysiology. Our data support previous findings that the serotonin, glutamate, hormonal and G-protein signalling pathways are involved in migraine. We suggest that a dys-function in neurotransmitter and hormone signalling is part of migraine pathophysiology.

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Competing interests
The authors report no competing interests.

Supplementary material
Supplementary material is available at Brain online.

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