Mutational Landscape of Autism Spectrum Disorder Brain Tissue

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Abstract: Rare post-zygotic mutations in the brain are now known to contribute to several neurodevelopmental disorders, including autism spectrum disorder (ASD). However, due to the limited availability of brain tissue, most studies rely on estimates of mosaicism from peripheral samples. In this study, we undertook whole exome sequencing on brain tissue from 26 ASD brain donors from the Harvard Brain Tissue Resource Center (HBTRC) and ascertained the presence of post-zygotic and germline mutations categorized as pathological, including those impacting known ASD-implicated genes. Although quantification did not reveal enrichment for post-zygotic mutations compared with the controls (n = 15), a small number of pathogenic, potentially ASD-implicated mutations were identified, notably in TRAK1 and CLSTN3. Furthermore, germline mutations were identified in the same tissue samples in several key ASD genes, including PTEN, SLC1A, CDH13, and CACNA1C. The establishment of tissue resources that are available to the scientific community will facilitate the discovery of new mutations for ASD and other neurodevelopmental disorders.

Keywords: exome sequencing; post-zygotic; somatic; germline; brain tissue; autism spectrum disorder (ASD)

1. Introduction

Autism spectrum disorder (ASD) is a complex neurodevelopmental disorder of early childhood onset, characterized principally by socio-communicative impairments and certain restricted behavioural patterns, but also associated with other neuropsychiatric and medical conditions. The genetic underpinning of this disorder is rapidly evolving, and the exact genetic mechanisms are complex and not fully understood, despite many ASD-implicated genes having been identified [1–4]. In brief, rare, particularly de novo, single nucleotide variants (SNV) and copy number variants (CNV) of variable size are implicated. These variants, whether SNVs or CNVs, are often variable in their penetrance and consequently almost certainly act in concert with each other and with other epigenetic and non-genetic factors. Moreover, genetic variants that are more commonly observed in the population (‘common variants’, minor allele frequency [MAF] ~1% or more) also play a role, but their discovery for ASD has lagged behind other neurodevelopmental and...
neuropsychiatric disorders [5]. The relative contribution of these different genetic factors and other non-genetic factors in any one individual is unclear, although cases have been described of Mendelian, or otherwise highly penetrant, single-gene/locus mutations [6,7].

Many mutations occur in the germline, inherited or arising de novo in one parent’s germ cell. However, post-zygotic mutations (PZM) may also occur during foetal development or, indeed, at any stage among those cells that undergo division as part of their life-cycle. Consequently, these events give rise to distinct cell populations, identifiable in peripheral samples, such as blood by a variant allele fraction that deviates from the 0.5 that would be expected for a typical heterozygous mutation. Using this approach, one recent study undertook ultra-deep sequencing of whole exomes from the Simons Simplex Collection and confirmed the contribution of genetic mosaicism to ASD [8]. Similarly, Lim and colleagues [9] investigated this using peripherally obtained DNA and determined that 7.5% of de novo mutations in ASD are post-zygotic, with particular genes enriched for these events.

For brain disorders, such mutations can also be directly identified in post-mortem brain tissue, a method that also allows a more detailed evaluation of their relationship to underly histopathological abnormality and brain expression [10,11]. Such an approach has been able to identify PZMs in resected brain tissue from patients with ASD [11,12] and epilepsy [13]. In one recent study using ultra-deep, whole-genome sequencing of 59 ASD donors, the enrichment of somatic SNVs was observed in neural enhancer sequences compared to 15 control donors [14]. This adds further weight to the evidence that post-zygotic mutations (PZMs) are important in ASD’s etiology and that they may shed light on specific pathophysiological mechanisms.

In order to further investigate the role of PZMs, we used post-mortem brain samples from the Harvard Brain Tissue Resource Center (HBTRC), obtained through the Autism Tissue Program, to investigate the presence of post-zygotic mutations in ASD donors. Although there are potential challenges to extracting DNA from brain tissue, we have previously demonstrated success [15], as well as shown how postmortem tissue can yield new, previously unidentified findings [16]. In our previous study, we examined these samples for CNVs and identified that several patients had CNVs of potential aetiological significance given their predicted deleterious impact and genomic context [15]. In this current study, deep exome sequencing was performed on these same post-mortem brain specimens from donors with ASD and on normal controls to identify mutations, including those we classify as likely to be post-zygotic.

2. Methods

Tissue preparation: The postmortem tissue samples were obtained from the Harvard Brain Tissue Resource Center (HBTRC). Ethical approval was obtained from The Hospital for Sick Children, REB number 0019980189. The cohort comprised 26 samples from ASD brain donors and 15 samples from controls. Among those classified as ASD, one is reported ‘suspected’ rather than confirmed and another is diagnostically labelled Angelman Syndrome. We previously examined 52 of these samples for the presence of CNVs [15]; to facilitate comparison, identical subject labels were used. The methods of DNA extraction are described in Wintle at al. [15] Briefly, the extraction of DNA from frozen samples was undertaken either manually, using the Gentra Puregene (Qiagen Inc., Germantown, MD, USA), or by using a semi-automated procedure (Gentra Autopure LS, Qiagen Inc., Germantown, MD, USA). Deep exome sequencing was performed to an average depth of 171X at the Centre for Applied Genomics (Toronto, ON, Canada) using established methods [17]. In brief, for short read sequencing, library preparation was performed from 500 ng of DNA using the TruSeq Nano DNA Library Preparation Kit, excluding the polymerase chain reaction (PCR) amplification step, followed by sequencing on an Illumina HiSeq X platform.

Variant Detection and Validation: Data analysis and base calling were performed using Bcl2FASTQ or HiSeq Analysis Software version 2.2.55.1311 (Illumina Inc., San Diego, CA, USA). Sequenced short reads were mapped to the hg19 reference sequence using the
Burrows-Wheeler Aligner [18], version 0.7.12. Single-nucleotide variations (SNVs) and indels were detected using Genome Analysis Toolkit (GATK), version 3.4-46. All the detected variants were annotated using ANNOVAR [19], which was setup with custom databases. All downstream quality control (QC) and bioinformatics analysis on the germline variants were performed using ANNOVAR annotated variants. To detect the post-zygotic variants, we used Mutect (Broad Institute, Cambridge, USA) [20]. The algorithm was run in pairs (proband-mother and proband-father) where the intersection of the two variant files was considered to be possible somatic variants in proband. We filtered out any variants that had an alternate variant ration <10% or >40%. This allowed us to detect high-level somatic mutations from our exome data.

We undertook many iterations of random validation across all variants using Sanger Sequencing and achieved a validation rate close to 100%. In this way, although all the prioritized variants were not individually validated, the quality of the data was supported.

Bioinformatic prediction of deleteriousness: We followed the American College of Medical Genetics (ACMG) guideline to assess variant clinical pathogenicity using ANNOVAR annotated vcf files. In brief, most of the common variants were assessed as benign. A rare variant (gnomAD frequency $< 5 \times 10^{-4}$) was assessed as clinically ‘pathogenic/likely pathogenic’ if it was listed as ‘pathogenic’ within the ClinVar database for other unrelated patients with similar phenotypes. There is scientific literature specifying the functional role of the gene in the disease physiology and multiple bioinformatics prediction tools to support the corresponding pathogenicity cutoff for the variant. A variant was assessed as a variant of uncertain significance (VOUS) if there was not much supported information available in the literature, but it was extremely rare in the control population. Deleterious mutations were defined according to three prediction tools: Polyphen2 [21], SIFT [22], and CADD [23]. Namely, all the variants with CADD $\geq 20$, and categorized as ‘damaging’ by SIFT and ‘probably damaging’ by Polyphen2, were defined as ‘bioinformatically deleterious’ by their corresponding cutoff values.

Gene set over-representation analysis: We explored for enrichment of post-zygotic genes using protocols implemented in both g:Profiler [24] and PANTHER [25]. Both methods examine over-representation of the gene list in each of a number of different gene-sets/pathways representing Gene Ontology (GO) annotation terms (in the case of PANTHER, this also includes PANTHER annotation terms) compared to a ‘universe’ of all genes in the genome. Fisher’s exact test was used to calculate a $p$ value, corrected for multiple comparisons with a Benjamini-Hochburg procedure.

3. Results

A total of 26 brain specimens from individuals with ASD (19 male, 7 female) and 15 from controls (12 male, 3 female) were included. Among the ASD cases, one was diagnosed with Angelman Syndrome and one with suspected ASD. The remaining 24 had all been diagnosed with ASD. All 26 cases are also reported in Wintle et al. [15], as are all 15 controls. Identical identification numbers are used across the two papers to facilitate comparison. One individual from Wintle et al. [15], who was diagnosed with epilepsy without ASD, did not have SNV data and is not included in the subsequent analyses. No further phenotype information is available on any of the cases or controls.

A total of 2766 SNVs were reported among the 26 cases (989 synonymous and 1777 non-synonymous) and 1485 among the 15 controls (504 synonymous and 981 non-synonymous). The mean number of variants per individual was 102 (synonymous: mean, SD = 36.1, 15.3; non-synonymous: mean, SD = 71.1, 21.6) for the cases and 99 (synonymous: mean, SD = 31.6, 5.5; non-synonymous: mean, SD = 65.4, 6.7) for the controls (Supplementary Spreadsheet).

3.1. Post-Zygotic Mutations

We were interested in identifying PZMs in brain tissue. By way of reminder, PZMs are those that arise at any stage after the formation of the zygote and give rise to distinct cell populations. This results in an allele fraction that deviates from the 0.5 that would
be expected for a heterozygous mutation. We used the established MuTect algorithm to
detect the PZMs. A total of 485 variants were classified as PZM, with 286 and 199 variants
in the cases and the controls, respectively (Supplementary Spreadsheet: post-zygotic—all;
post-zygotic—predicted damaging). Two hundred and seventy-one of these variants were
non-synonymous (159 in the cases and 112 in the controls), of which 25 were categorized as
pathogenic according to the bioinformatic prediction (CADD, SIFT, and PolyPhen), 11 in
11 different controls and 14 in 11 cases. Of these, one was deemed likely to be pathogenic
according to the ACMG guidelines [26] (Table 1). No enrichment for damaging PZMs was
therefore observed in the cases versus the controls (Fisher’s Exact Test, ns). None of these
variants was observed in the data from the 1000 Genomes Project, and all were either not
observed or observed with extremely low frequency ($<5 \times 10^{-4}$) in GnomAD.

None of the 14 genes with bioinformatically predicted damaging mutations overlapped
the ‘ASD-implicated’ genes, a curated list of nearly 1000 putative ASD genes with varying
degrees of supporting evidence [2]. Moreover, none of the genes were observed in the list of genes
with high confidence of the ASD somatic mutations identified in Lim et al. [9]. Although
none of the genes was previously implicated in ASD [27], one, TRAK1 [chr3:42,244,127
(hg19), A > G], interacts with the known schizophrenia gene DISC1 [28,29], and one other
gene, CLSTN3 [chr12:7310,284 (hg19), G > A], was deemed likely to be pathogenic according
to the ACMG guidelines. This gene encodes a calsyntenin, proteins which as a group are
known to function as synaptogenic adhesion molecules in concert with others such as the
neurexins, themselves implicated in ASD and other neurodevelopmental disorders [30].
Taken together with the predicted damaging nature of this mutation, it is quite possible
that this was aetiollogically related to the ASD diagnosed in this individual. Unfortunately,
no additional clinical information is available from this subject.

We next examined those genes impacted by predicted damaging variants among cases
for over-representation of different curated gene sets, as implemented in PANTHER and
g:profiler. No over-representation in any of the examined gene sets was observed. We also
examined protein-protein interaction pathways in STRING, but no pathway enrichment
was observed. We also compared the case and control genes for any enrichment in brain
expression at particular developmental time points (prenatal, early childhood, and adult in
the cases versus the controls), but no differences were observed (Supplementary Table S1).

### 3.2. Germline Mutations

A total of 3749 germline mutations were identified, with 2460 in the cases and 1288 in
the controls. Among those with ASD, there were 1618 non-synonymous variants, among
which 200 were bioinformatically predicted as deleterious, including seven stop-gain
mutations (Supplementary Spreadsheet: germline—all; germline—predicted damaging).
None of these variants was observed in the 1000 Genomes dataset, and all were either not
observed or occurred with extremely low frequency ($<5 \times 10^{-4}$) in GnomAD. Thirteen
germline mutations were deemed pathogenic or clinically pathogenic according to the
ACMG guidelines (Table 1).

Seventy of the genes impacted by the 200 bioinformatically predicted damaging
mutations are reported in Lim et al. (2017). [9] Eight of the genes impacted by these
200 variants are in the ASD-implicated list of genes (Table 1). Among these are one
stop-gain mutation in PTEN [chr10:89,685,300 (hg19), C > A] and a missense mutation in
SCN1A [chr2:166,856,252 (hg19), G > A]. Both PTEN [31] and SCN1A [32,33] are established
neurodevelopmental genes, with mutations recorded in association with several disorders,
including ID, epilepsy, and ASD. In view of their established implication in ASD and other
neurodevelopmental disorders, we hypothesize that these mutations played an etiological
role in the ASD diagnoses in both cases.
Table 1. Characteristics of rare, predicted damaging mutations clinically classified as pathogenic or likely pathogenic, identified in brain tissue.

| Sample | Type | LocusID | Ref | Alt | GnomAD | Gene | Classification | Band | Pos | Size | Type | Genes |
|--------|------|---------|-----|-----|--------|------|----------------|------|-----|------|------|-------|
| AN00090 | -    | -       | -   | -   | -      | -    | -              | -    | -   | -    | Loss | Many  |
| AN01093 | -    | -       | -   | -   | -      | -    | -              | 7q34 | 142,538,076–142,561,946 | 23.9 | Loss | PIP   |
| AN01570 | Germline chr7:1510817:1510817 | C T | 8.4 × 10^{-6} | INTS1 | Pathogenic | -    | -              | -    | -   | -    | -    | -     |
| AN03345 | Germline chr6:43008300:43008300 | G C | NO  | PTEN | Pathogenic | -    | -              | -    | -   | -    | -    | -     |
| AN03935 | Germline chr12:23687394:23687394 | G T | NO  | SOX5 | Pathogenic | 15q11.1-q13.2 | 18,278,739–28,280,653 | 10,001.90 | Gain | Many  |
| AN06420 | Germline chr13:23904448:23904448 | C T | NO  | SAC5 | Likely pathogenic | 6q26 | 162,863,051–162,917,072 | 54 | Loss | PARK2 |
| AN07664 | Germline chr8:133187845:133187845 | G A | NO  | KCNQ3 | Pathogenic | 19p13.42 | 60,601,790–60,943,899 | 342.1 | Gain | Many  |
| AN08043 | Germline chr16:83711952:83711952 | C T | NO  | CDH13 | Pathogenic | -    | -              | -    | -   | -    | -    | -     |
| AN08166 | Germline chr2:8871659:8871659 | A G | 4.9 × 10^{-5} | KIDINS220 | Likely pathogenic | -    | -              | -    | -   | -    | -    | -     |
| AN08873 | Germline chr12:7310284:7310284 | G A | NO  | CLSTN3 | Likely pathogenic | -    | -              | -    | -   | -    | -    | -     |
| AN09402 | Germline chr3:158383151:158383151 | G A | 8.1 × 10^{-6} | GFM1 | Likely pathogenic | 15q11.1-q13.1 | 18,276,341–26,752,537 | 8476.20 | Gain | Many  |
| AN1049    | -    | -       | -   | -   | -      | -    | -              | 8q22.1 | 95,265,603–95,304,988 | 39.4 | Gain | CDH17 |
| AN13872   | -    | -       | -   | -   | -      | -    | -              | 16p13.2 | 6,992,775–7,021,963 | 29.2 | Loss | A2BP1 |
| AN14613   | -    | -       | -   | -   | -      | -    | -              | 3p14.2 | 60,464,015–60,502,990 | 39 | Loss | FHT   |
| AN14762   | -    | -       | -   | -   | -      | -    | -              | 2p16.3 | 51,075,080–51,087,539 | 12.5 | Loss | NRXN1 |
| AN14829   | Germline chr19:42857121:42857121 | C T | 8.2 × 10^{-6} | MEGF8 | Likely pathogenic | Xp22.33 | 2,281,299–2,493,943 | 212.6 | Gain | ZBED1, DHRX |
| AN16115   | Germline chr2:166856252:166856252 | G A | NO  | SCN1A | Pathogenic | 15q11.1-q13.2 | 18,276,341–28,289,587 | 10,013.20 | Gain | Many  |
| AN16641   | Germline chr12:2717736:2717736 | G A | NO  | CACNA1C | Pathogenic | -    | -              | -    | -   | -    | -    | -     |
| AN17138   | -    | -       | -   | -   | -      | -    | -              | 7p21.1 | 16,315,368–16,370,511 | 55.1 | Loss | hCG_1745121/ISPD |
| AN19511   | -    | -       | -   | -   | -      | -    | -              | 15q11.2 | 20,302,458–21,937,715 | 1635.30 | Gain | Many  |
| AN19511   | -    | -       | -   | -   | -      | -    | -              | 15q11.2 | 21,985,041–22,943,182 | 958.1 | Gain | Many  |
| AN19511   | -    | -       | -   | -   | -      | -    | -              | 15q11.2 | 22,989,278–23,915,837 | 926.6 | Gain | Many  |
| AN19511   | -    | -       | -   | -   | -      | -    | -              | 15q12-q13.1 | 23,925,463–26,500,067 | 2574.60 | Gain | Many  |
| AN19511   | -    | -       | -   | -   | -      | -    | -              | 4p15.31 | 22,351,359–22,429,602 | 78.2 | Loss | GBA3   |

1 Frequency of variant in GnomAD database.
We also identified variants impacting CDH13 [chr16:83,711,952 (hg19), C > T] and CACNA1C [chr12:2717,736 (hg19), G > A]. CDH13 is a Cadherin gene, a group of genes which encode a class of calcium-dependent transmembrane protein; variants in this gene have been implicated in neurodevelopmental phenotypes, including ASD and ADHD [34]. CACNA1C is another calcium channel protein. Common variants in this gene have been identified in association with several different neuropsychiatric phenotypes [35,36], including ASD [37]. It is quite possible, therefore, that these mutations also played an etiological role in the ASD diagnosis in these cases.

Five samples were identified as having both a rare, predicted damaging single nucleotide variant and one or more pathogenic CNVs, raising the possibility of two potentially causative genetic mutations. For example, one sample had a likely pathogenic germline variant impacting GFM1 and a CNV deletion at the MACROD2 gene. Another sample had both a predicted pathogenic SNV in GRM1 and a CNV loss impacting PARK2.

We studied the over-representation of the genes impacted by the predicted damaging variants in the cases in the curated gene sets, as described above. No over-representation was observed for either the PZM or the germline genes in any of the gene sets studied. Additionally, no enrichment in the cases versus the controls was observed for genes with expression across the prenatal, early childhood, or adult developmental stages (Supplementary Table S1).

4. Discussion

Brain specimens are a rare resource in neurodevelopmental disorders but can uniquely provide insight into the genetic mechanisms of disease through the ability to directly study the very tissues that underlie the manifestations of these disorders [9–13,15]. This includes the ability to identify the tissue of specific post-zygotic mutations. Such mutations have for a long time been recognized as an important etiological factor in focal cortical dysplasias and epilepsy syndromes [13]. However, more recently, their role in other neurodevelopmental disorders, including ASD, has emerged [11,12,15]. Much of the evidence is based on the detection of alternate allele fraction (AAF) from the deep sequenced coverage of peripherally derived specimens. In the current study, we examined donated brain tissue from patients with ASD and identified PZMs in several genes that may be aetiologically implicated in their ASD diagnoses. Although none of the genes harbouring predicted-damaging PZMs was previously implicated in ASD, intellectual disability (ID), or other neurodevelopmental disorders, two, TRAK1 and CLSTN3, are widely expressed in the brain.

TRAK1 is closely associated with DISC1, itself implicated in schizophrenia and neural development [29,38]. Mutations in DISC1 have been well documented in association with schizophrenia but not ASD. Additionally, one predicted pathogenic mutation in TRAK1 has also been described in the literature in a subject with schizophrenia [39]. The implications, therefore, for neurodevelopmental disorders such as ASD are unclear, although the genetic relationship between ASD and schizophrenia is now well-established [40]. Stronger evidence for a possible role in ASD phenotypes exists for CLSTN3, a calsyntenin, which as a group function as synaptogenic adhesion molecules [30,41]. Specifically, CLSTN3 is thought to be important for the normal development and function of the GABAergic and glutaminergic synapses, and although not specifically identified as an ASD-implicated gene, a de novo predicted damaging mutation in this gene has previously been described in an ASD proband [42]. Rodent models of CLSTN3 have identified the impact of deletion on impairing cognitive function [43], offering a possible pathophysiological mechanism.

In contrast to our search among the PZMs, we did identify several germline mutations that are likely implicated in the ASD diagnoses of these subjects. For example, a mutation in PTEN was identified in one subject. PTEN is an overgrowth gene, with a variable phenotype that may include macrocephaly, tissue proliferation, including tumors, and neurodevelopmental disorders [31]. Indeed, research suggests that as many as 20% of individuals with autism who also are macrocephalic may have a PTEN mutation [44].
another subject, a predicted damaging mutation in \textit{SCN1A} was identified. \textit{SCN1A} encodes
the $\alpha$-1 subunit of the NaV1.1 sodium channel and is strongly associated with epilepsy as
one of the most important channelopathies \cite{32,33}. Its most notable association is Dravet
Syndrome, but other epilepsy and movement disorders have also been described, as has
ASD, although the exact prevalence is unclear, and may be in the region of 24–40\% \cite{45}. We
do not know if individual AN16115 had epilepsy during their lifetime.

We also identified variants impacting \textit{CDH13} and \textit{CACNA1C}. \textit{CDH13} is a Cadherin
gene, a group of genes which encode a class of calcium-dependent transmembrane protein.
Their embryonic expression earmarks the critical role they play in axonal growth and
synapse development. Although there are many members of this protein superfamily, a
small number have been described in association with neurodevelopmental phenotypes,
including ASD and ADHD \cite{34}. \textit{CACNA1C} is another calcium channel protein, encod-
ing the $\alpha$-1c subunit of the L-type voltage-gated calcium channel. Common variants
in this gene have been identified in association with several different neuropsychiatric
phenotypes \cite{35,36}, including ASD \cite{37}. Moreover, mouse models have suggested a role
for this gene in potentially related cognitive endophenotypes that include fear condition-
ing \cite{46}. It is quite possible, therefore, that these mutations also played an etiological role
in the ASD diagnosis in these cases and further reinforce the importance of cadherins and a
channelopathy-mediated pathogenesis in ASD.

This current study was conducted to show the potential utility of studying patterns of
mutation in brain tissue and specifically to identify post-zygotic mutations that may be of
aetiological relevance. Although we failed to identify predicted damaging mutations in any
known ASD or other neurodevelopmental genes, our study has successfully identified post-
zygotic mutations, including a small number in genes that warrant further investigation.
Our results may alternatively suggest a less widespread role for PZMs than previously
predicted. However, considering the emerging evidence from epilepsy and ASD discussed
above, this seems unlikely. The small sample size of the current study will have impacted
on the power of this study to identify these variants. It is also possible that PZMs are of
aetiological significance only for particular phenotypes, such as severe ASD, nonverbal ASD,
or ASD with co-morbidities such as epilepsy. In the absence of phenotype information, we
are unable to further comment on this; however, moving forward, any brain tissue resource
will need to also include detailed phenotype information to facilitate the unravelling
of these brain-behaviour connections. Moreover, we are not able to indicate whether
undiected brain regions in these specimens are also negative for PZMs. As discussed in
Wintle et al. \cite{15}), the BA17 visual cortex was used for this current study, in part due to
its easy dissection and availability. Ideally, the search for PZMs will be predicated on the
identification of histopathological abnormality, which will then motivate the decision on
which regions of the brain to dissect.

We hope that our results will additionally provide a resource of genetic results on
postmortem tissue in ASD patients. The nature of these mutations, including their mecha-
nism, the genes impacted, and the potential effects on brain development will only be truly
resolved with greater emphasis on studies such as this one that are also able to evaluate
histology and cellular function in the tissues impacted. There are several ways in which
studies such as this can make further progress. The establishment of tissue resources that
are available to the scientific community are making samples available, allowing increased
sample sizes, and these are often from patients who have detailed phenotype information.
In considering sample size, there are a number of operational complexities in establishing
a tissue resource. Samples may have to be collected from different geographic locations,
and these may have been stored under different conditions. Detailed information will be
needed on factors that may impact the results and their interpretation. Moreover, whereas
on the one hand the tissue may comprise samples resected during surgery, for other families
the tissue may have been donated after the death of a loved one. This requires a sensitive
and ethical process to be in place. Consequently, we anticipate that sample sizes will
remain modest in comparison to other methodological approaches. This notwithstanding,
the strength of this approach is the ability to directly identify PZMs in brain tissue as well as more closely study the impact of mutations in the tissue itself. In the future, it will be important to more carefully examine genetic architecture across different regions; this will be facilitated by using single cell genomic approaches. Additionally, it will be important to stratify samples by sex and developmental age, as well as to compare the genetic architecture between histologically normal and abnormal tissues.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/genes13020207/s1, supplementary spreadsheet comprising all identified variants across samples. Table S1: Examination of expression profiles of predicted damaging mutations.

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Conflicts of Interest: S.W.S. is on the Scientific Advisory Committees of Deep Genomics, Population Bio, and is an Academic Consultant for the King Abdulaziz University. The authors declare no conflict of interest.

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