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TITLE: MTHFR Functional Polymorphism C677T and Genomic Instability in the Etiology of Idiopathic Autism in Simplex Families

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14. ABSTRACT
Autism Spectrum Disorder (ASD) are a group of neurodevelopmental disorders that are caused by a range of factors, including: genetic, epigenetic and environmental, with a genetic/epigenetic model proposed (Jiang et al., 2004). While a main focus of autism research remains on the genetic causes, more and more attention was drawn to the role epigenetic factors play, as it has been shown to play a role in idiopathic autism. With our previous published study revealed significantly association of C677T polymorphism in MTHFR gene with idiopathic autism in Simplex (SPX) autism families (Liu et al., 2011); and the proven facts that de novo CNVs rates are consistently high in SPX ASD (5.8% - 10.2%) versus familial ASD (2 - 3%), we hypothesize that low-activity MTHFR 677T allele leads to increase global DNA hypomethylation and consequently results in increased generation of de novo CNVs bringing about a higher risk for developing sporadic cases of autism. We proposed to test 1) the association of MTHFR 677T allele with rate of ASD related de novo CNVs; 2) the association of of MTHFR 677T allele with increased level of global hypomethylation; and 3) the association of level of global hypomethylation with increased rate of ASD related de novo CNVs.

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INTRODUCTION:

Autism Spectrum Disorder (ASD) are a group of neurodevelopmental disorders that are caused by a range of factors, including: genetic, epigenetic and environmental, with a genetic/epigenetic model proposed (Jiang et al., 2004). While a main focus of autism research remains on the genetic causes, more and more attention was shawn to the role epigenetic factors play, as it has been shown to play a role in idiopathic autism. With our previous published study revealed significantly association of C677T polymorphism in MTHFR gene with idiopathic autism in Simplex (SPX) autism families (Liu et al., 2011); and the proven facts that de novo CNVs rates are consistently high in SPX ASD (5.8%-10.2%) versus familial ASD (2-3%), we hypothesize that low-activity MTHFR 677T allele leads to increase global DNA hypomethylation and consequently results in increased generation of de novo CNVs bringing about a higher risk for developing sporadic cases of autism. We proposed to test 1) the association of MTHFR 677T allele with rate of ASD related de novo CNVs; 2) the association of of MTHFR 677T allele with increased level of global hypomethylation; and 3) the association of level of global hypomethylation with increased rate of ASD related de novo CNVs.

KEYWORDS

Autism, Sporadic Cases, MTHFR, Hypomethylation, Differentially Methylated Regions (DMR), Copy Number Variation (CNV)

OVERALL PROJECT SUMMARY

This pilot project started from September of 2013. During the over two years of project execution, we have achieved the aims set in the original proposal. Although no publication has yet come out from the study during till today, the significant findings from the study allow us to prepare a significant manuscript that we are aim at submitting to a top journal during this March; and a follow-up study which was carried out with PI's other funding source will very likely yield another manuscript during the middle of this year.

KEY RESEARCH ACCOMPLISHMENTS:

- Completed microarray data analysis on 510 SPX families; identified 99 individuals carrying pathogenic CNVs; confirmed 33 of these CNVs be de novo.
- 510 SPX families (both parents and affected individuals) were genotyped on MTHFR Functional Polymorphism C677T using TaqMan Assay.
- Both experimental and bioinformatics pipelines were established for global Methylation profiling using MBD-Seq strategy with Ion Torrent Proton
- 31 ASD cases with de novo pathogenic CNVs and 31 ASD cases without de novo pathogenic CNVs were MBD-Sequenced. Among all 62 ASD cases, at MTHFR C677T locus, 35, 23 and 4 carry CC, CT and TT genotype respectively.
- The following data analyses were performed
  1. **The MBD-Seq data QA/QC:** Data quality was measured in a number of ways to ensure that the data was of good quality prior to analysis. The first was a measure of saturation, with a cutoff minimum score of 0.5, to determine if the data creates reproducible coverage of the reference genome. The second measure was enrichment, with a minimum cutoff score of 1.7, which measures how well the methylation capture method worked and therefore how enriched the sample is for methylated DNA. The third measure was 5 times or greater coverage, with a minimum cut-off of 5%, meaning that at least 5% of the sequence needed to have at least 5 times sequencing coverage. The fourth measure was the number of total sequencing reads for a sample, with a minimum of 20 million reads. The fifth measure was the number of unique reads for a sample, with a minimum of 15
million reads as a cutoff, where a unique read was defined as a read that does not share a stop or start point with any other read. The number of reads, both total and unique, were plotted to visualize all samples individually to ensure they met this criteria. **Overall, all 62 samples passed all five quality measurements and, thus, were included in the analysis.**

2. **Global Methylation Index (GMI):** The global methylation index (GMI) was calculated across the genome for each sample. The average GMI for each intended comparison group, such as group carrying CNV and group not carrying CNV; group carrying genotype C/C and group carrying C/T and T/T genotype on MTHFR C677T polymorphism, was calculated, were compared between groups using t-tests. While no significant difference was found in global methylation level between CNV+ and CNV- groups, we did find the low activity T allele ASD carrier for MTHFR C677T variant has a significantly lower global methylation level than the CC homozygous group.

3. **Detection of Differentially Methylated Regions (DMRs) between C/C and C/T+T/T groups:** The methylation level within a 200bp window was calculated and normalized to reads per kilobase per million (RPKM). Following a comparison of RPKM values in each 200 bp window between C/C and C/T+T/T groups at p=0.05, 238 differentially methylated regions (DMRs) were identified. 140 DMRs were hypermethylated in the C/C group relative to the C/T+T/T group. 98 DMRs were hypermethylated in the C/T+T/T group relative to the C/C group. The RPKM values for the DMRs that were found to be hypermethylated in the C/T+T/T group were, generally, lower overall when compared to those identified as hypermethylated in the C/C group. **It is of great interest that the identified DMRs were not evenly distributed across chromosomes.**

4. **Most Significantly:** The differentially methylated regions (DMRs) were found to be highly biased towards autism related genes and CpG islands. This may imply a major mechanism for etiology of sporadic cases of autism: autism causing environmental factors serves as modulator to regulate a MTHFR -mediated epigenomics to regulate specific autism related gene.

**CONCLUSION:**

The proposed research aims for the funded pilot projects were successfully realized with a significant finding for a potential major mechanism for etiology of sporadic cases of autism.

**PUBLICATIONS, ABSTRACTS AND PRESENTATIONS:** N/A

**INVENTIONS, PATENTS AND LICENSES:** N/A

**REPORTABLE OUTCOMES:**

The significant findings from the study allow us to prepare a significant manuscript that we are aim at submitting to a top journal during this March; and a follow-up study which was carried out with PI's other funding source will very likely yield another manuscript during the middle of this year.

**OTHER ACHIEVEMENTS:** N/A

**REFERENCES:**
**MBD-Seq Data Quality**

Data quality was measured in a number of ways to ensure that the data was of good quality prior to analysis. The results of these measurements for each sample are summarized in Table 1. The first was a measure of saturation, with a cutoff minimum score of 0.5, to determine if the data creates reproducible coverage of the reference genome (Lienhard *et al.*, 2014). The second measure was enrichment, with a minimum cutoff score of 1.7, which measures how well the methylation capture method worked and therefore how enriched the sample is for methylated DNA. The third measure was 5 times or greater coverage, with a minimum cutoff of 5%, meaning that at least 5% of the sequence needed to have at least 5 times sequencing coverage. Figure 1 is an example of data quality output, while Figure 2 demonstrates the distributions of the quality metrics for each sample group.

**Figure 1: Example Quality Control Graphs of Subject 21.**

Graphs created during MeDIPs data quality control. A: saturation analysis determines if the data generated creates a reproducible coverage of the reference genome (Lienhard *et al.*, 2014); B: representation of amount of times (X) coverage achieved with data generated as a percentage of all data generated. Total >=5X coverage is the summation percentage of 5-5X and >5X coverage.
Figure 2: Data Quality Metrics Grouped by Genotype

Boxplots of quality control measures of the data grouped by rs1801133 (C677T) genotype groups. Each sample data needed to pass all three thresholds to be considered for analysis. A: percentage of at least 5 times coverage (minimum = 5%); B: relative methylation enrichment score (minimum = 1.7); C: saturation score (minimum = 0.5).

The fourth measure was the number of total sequencing reads for a sample, with a minimum of 20 million reads. The fifth measure was the number of unique reads for a sample, with a minimum of 15 million reads as a cutoff, where a unique read was defined as a read that does not share a stop or start point with any other read. The number of reads, both total and unique, were plotted to visualize all samples individually to ensure they met this criteria; this can be seen in Figure 3.

Overall, all 62 samples passed all five quality measurements and, thus, were included in the analysis.
At least 20 mil reads were needed, with at least 15 mil of those reads being classified as unique, to pass quality control.

**Global Methylation Index (GMI)**

The global methylation index (GMI) was calculated across the genome for each sample. The average GMI for each genotype group (C/C, C/T and T/T) was calculated, were compared using t-tests. Following this preliminary analysis, it was decided that the C/T and T/T groups could be combined into one group for further analysis because there was no significant difference in mean GMI between the C/T and T/T (data not shown).

The mean GMI for C/C genotype group was 47.22, with a standard deviation of 18.41 and n=35. The mean GMI for C/T+T/T genotype group was 40.22, with a standard deviation of 9.37 and n=27. Comparing the mean GMI between C/C genotype group and the C/T+T/T genotype group using a t-test showed that there was a marginally statistically significant difference between the groups ($p=0.0569$; Figure 4).
Samples were grouped into two groups: C/C genotype and genotypes with a T present (C/T and T/T). A two-tailed t-test, with independent samples and unequal variances was run using IBM SPSS Statistics v22 to compare the mean global methylation index (GMI), where n=35 for C/C and n=27 for C/T+T/T.

*: significantly different, p=0.0569.

Analysis of Differentially Methylated Regions (DMRs)

The methylation level within a 200bp window was calculated and normalized to reads per kilobase per million (RPKM). Following a comparison of RPKM values in each 200 bp window between C/C and C/T+T/T groups at p=0.05, 238 differentially methylated regions (DMRs) were identified. These identified DMRs are summarized in a table not presented here. 140 DMRs were hypermethylated in the C/C group relative to the C/T+T/T group. 98 DMRs were hypermethylated in the C/T+T/T group relative to the C/C group. The RPKM values for the DMRs that were found to be hypermethylated in the C/T+T/T group were, generally, lower overall when compared to those identified as hypermethylated in the C/C group (see Figure 5).
Figure 5: Reads per Kilobase per Million (RPKM) Mean Comparison Between Genotype Groups Across Differentially Methylated Regions (DMRs)

Comparison of mean RPKM values across DMRs when C/T + T/T genotype group was hypermethylated (blue dots) and when C/C genotype group mean was hypermethylated (orange dots).

We found that the identified DMRs were not evenly distributed across chromosomes. This phenomenon was quantified by comparing the actual number of DMRs per chromosome identified to a calculated number of DMRs expected for each chromosome based on chromosome size. It was found that 16 of the 24 chromosomes had a significantly different actual number of DMRs compared to the expected number of DMRs (see Figure 7). Six of the chromosomes were found to be significantly enriched for DMRs while 10 of the chromosomes were found to be reduced in DMRs.
Figure 7: Genomic Distribution of Differentially Methyalted Regions (DMRs)

The actual number of DMRs for each chromosome at p=0.05 versus the number of DMRs expected per chromosome if the DMRs were distributed evenly based on chromosome size (hg19/GRCh37 Feb 2009; http://www.ncbi.nlm.nih.gov/projects/assembly/human/data/). There was no data available for mitochondrial DNA.

*: significantly different from expected, p<0.05

Sequenom Validation

To validate the findings by MBD-Seq, an alternate method – Sequenom MassARRAY – was chosen to analyze a selection of the DMRs identified. Of the 59 DMRs identified in intragenic regions, 8 of these were selected for validation. Six of the DMRs were hypermethylated in the C/C group and two were hypermethylated in the C/T+T/T group. The data is under analysis.