Genetic Analysis of Neuroligin 4Y Gene in Autism Population of India

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Abstract

Background Autism is one of the most complex, heterogeneous neurological disorders. It is characterized mainly by abnormal communication, impaired social interaction, and restricted behaviors. Prevalence of autism is not clear in Indian population.

Aim The present study hypothesized that Y chromosome plays role in sex bias of autism in Indian autistic population. To investigate our hypothesis, we underwent genetic analysis of neuroligin 4Y [NLGN4Y] gene by sequencing 85 male autistic children after screening large population of 1,870 mentally ill children from North Karnataka region of India.

Result Detailed sequencing of the single targeted gene revealed nine variants including, one novel missense mutation and eight synonymous variants; this accounts for 88.9% of synonymous variants. A single novel missense mutation is predicted to be nonpathogenic on the functions of neuroligin4Y protein but it slightly affects the local configuration by altering the original structure of a protein by changing charge and size of amino acid.

Conclusion Probably NLGN4Y gene may not be the risk factor for autism in male children in Indian autistic population. Functional analysis was an important limitation of our study. Therefore, detailed functional analysis is necessary to determine the exact role of novel missense mutation of neuroligin 4Y [NLGN4Y] gene especially in the male predominance of autism in Indian autistic population.

Keywords
► autism
► neuroligin 4Y
► India
► novel missense mutation
► male predominance

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Introduction

Autism [ASD] [MIM 299850] is a heterogeneous neurodevelopmental disorder. Autism is not characterized based on a single symptom. It is usually characterized by the triad of symptoms viz lack of social interaction, abnormal verbal and nonverbal communications and stereotyped or repetitive behaviors.\(^1\) Autism is classified as syndromic and non-syndromic autism. Syndromic autism is one in which patients who have pre-existing neurological disorders, example, a subset of patients with fragile x syndrome, tubers sclerosis, Rett syndrome displays phenotypes which are attributed to ASD. Non-syndromic autism accounts for autism cases that are not linked to any neurological disorders.\(^2\) Neuroligin is trans-synaptic cell adhesion molecule present post-synaptically and plays a very important role in synaptogenesis with presynaptic neurexin.\(^3\) Humans have neuroligin 4X [NLGN4X] on the X chromosome and neuroligin 4Y [NLGN4Y] on the Y chromosome. NLGN4X and NLGN4Y genes share 97% sequence identity.\(^4\) The male bias seen from NLGN4X mutations is unclear since NLGN4Y plays a function similar to NLGN4X and should be sufficient to reimburse for NLGN4X ASD-related mutations. This lack of compensation in males suggested that NLGN4Y may have an uncharacterized distinct function that needs to be explored. Several studies are reported that synaptic cell adhesion molecules have been strongly involved in autism. Neuroligin has an important role in the maturation and functions of synapses.\(^5,6\) The mechanism of Y chromosome contribution on to neurodevelopmental disorders is still not known very well. Originally, it was thought that Y chromosome contains only a few genes that are primarily involved in sex determination and testicular functions but now it is known to contain numerous genes with diverse functions.\(^7\) Several shreds of evidence strongly suggested that NLGN4X deficiencies can cause autism and still there is no clear understanding of sex bias in autism.

We hypothesize that male individuals have both X and Y chromosome so analysis of sequence variants in NLGN4Y gene may be associated with sex bias in male autistic individuals. To address this objective, we sequenced all the exonic regions of NLGN4Y gene in 85 male autistic children from north Karnataka region of India.

Methods

Sample Collection

One-hundred fifty autistic children were identified after screening a large mentally ill population of 1,870 children from the entire North Karnataka region of India \((n_{\text{male}} = 117, n_{\text{female}} = 33, \text{mean age} = \pm 11.5)\) which accounts for 8.02% of autism in North Karnataka region of India. Sanger Sequence analysis of neuroligin 4Y gene from 85 male autistic children revealed the nine variants, which include one missense and eight synonymous variants. Four variants which were recorded in our study cohort are not previously recorded in any in house human SNP databases viz dbSNP, 1000 genomes, ExAc and ClinVar shown in \(\text{Table 2}\). Novel missense, p.N163K mutation was recorded in three autistic children and clinical features of those autistic children with missense mutation are shown in \(\text{Table 1}\). Pathogenicity prediction of missense variants was analyzed using Insilco tools like PROVEAN, SNAP2, polyphen2, SNP&GO, and CADD. Conservation status analysis of amino acid residues of NLGN4Y protein at 163 position was checked using the Consurf Server (https://consurf.tau.ac.il/).\(^8\) Three-dimensional structure of wild type and mutant protein was developed using Swissmodel and structures were visualized and analyzed using UCSF Chimera tool.

Results

Detailed screening of 1,870 mentally ill children below 18 years of age from North Karnataka population of India revealed 150 autistic children \((n_{\text{male}} = 117, n_{\text{female}} = 33, \text{mean age} = \pm 11.5)\) which accounts for 8.02% of autism in North Karnataka region of India. Sanger Sequence analysis of neuroligin 4Y gene from 85 male autistic children revealed the nine variants, which include one missense and eight synonymous variants. Four variants which were recorded in our study cohort were not previously recorded in any in house human SNP databases viz dbSNP, 1000 genomes, ExAc and ClinVar shown in \(\text{Table 2}\). Pathogenicity prediction of missense variants was analyzed using Insilco tools viz PROVEAN, POLYPHEN2, SNAP2, and CADD. Only Missense variant, p.N163K was found to be harmless on the functions of NLGN4Y protein by PROVEAN, POLYPHEN2, SNAP2, SNP&GO, and CADD shown in \(\text{Table 3}\). Conservation status analysis of NLGN4Y protein sequences shows that amino acid residue at 163 position is not conserved; it is variable and exposed residue according to the neural network algorithm shown in \(\text{Fig. 1}\).

Three-dimensional protein modeling analysis of NLGN 4Y protein revealed that mutant residue is bigger than wild type.
residue and it possesses a positive charge whereas wild type protein possesses neutral charge. The wild type of residue is predicted to be located in its preferred secondary structure, a turn but the mutant residue prefers to be in another secondary structure; therefore, the local conformation will be slightly destabilized shown in \(\text{Fig. 2A}\) and \(\text{Fig. 2B}\). The mutated residue is located in a domain that is important for binding of other molecules. Mutation of the residue might disturb this function of NLGN4Y protein.

### Discussion

Autism [MIM 299850] is a complex neurological condition which is characterized by abnormal social interaction, verbal and nonverbal communication and impaired behaviors. “Autism,” the term was first used by Ukrainian-Austrian-American psychiatrist, Leo Kenner in 1943.\textsuperscript{9} Rates of autism cases are increasing globally over the period of time when it comes to the Indian perspectives, cases are increasing dramatically, and it may be due to increased scientific knowledge and awareness or it may be an improper diagnosis. In recent days, neuroligin gene is the most targeted gene for the molecular studies on neurological disorders like autism, anxiety, attention deficit hyperactivity disorder and intellectual disability due to its active role in synaptogenesis.\textsuperscript{10–13}

The Simons Foundation Autism Research Initiative (SFARI) [geneSFARI.org] lists four genes of Y chromosome associated with autism viz NLGN4Y, ASMT, USP9Y, and SHOX.\textsuperscript{14} Only a few studies have been undertaken till now to study the role of NLGN4Y gene in autism. Studies conducted in 2005 and 2006, failed to identify the variants in the NLGN4Y gene in autistic patients.\textsuperscript{15,16} But later several studies, record polymorphisms

### Table 1 Showing list of mutations recorded in our study cohort

<table>
<thead>
<tr>
<th>Variation</th>
<th>N. change</th>
<th>A.A Change</th>
<th>SNP id</th>
<th>Frequency of mutation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Missense</td>
<td>g. 205526 C &gt; A</td>
<td>p.N163K</td>
<td>Not recorded</td>
<td>3 (3.5%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312652 T &gt; C</td>
<td>p. H447=</td>
<td>rs777234513</td>
<td>4 homozygous (4.7%) 2 heterozygous (2.3%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312781C &gt; T</td>
<td>p.G490=</td>
<td>rs767447455</td>
<td>3 (3.5%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312787 A &gt; G</td>
<td>p.E492=</td>
<td>rs750273940</td>
<td>3 homozygous (3.5%) 2 heterozygous (2.3%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312826 A &gt; C</td>
<td>p.T505=</td>
<td>Not recorded</td>
<td>1 (1.2%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312844 T &gt; C</td>
<td>p.N512=</td>
<td>Not recorded</td>
<td>2 (2.3%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312847 C &gt; T</td>
<td>p.F513=</td>
<td>Not recorded</td>
<td>1 (1.2%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312871 A &gt; C</td>
<td>p.S520=</td>
<td>rs1423308667</td>
<td>1 (1.2%)</td>
</tr>
<tr>
<td>Synonymous</td>
<td>g.312880 G &gt; C</td>
<td>p.V523=</td>
<td>rs753006927</td>
<td>3 (3.5%)</td>
</tr>
</tbody>
</table>

### Table 2 Clinical features of autism children with missense mutation, p.N163K of NLGN4Y gene

<table>
<thead>
<tr>
<th>Demographic character</th>
<th>Child 1</th>
<th>Child 2</th>
<th>Child 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethnic origin</td>
<td>Indian</td>
<td>Indian</td>
<td>Indian</td>
</tr>
<tr>
<td>Sex</td>
<td>Male</td>
<td>Male</td>
<td>Male</td>
</tr>
<tr>
<td>Age of father at child’s birth</td>
<td>38</td>
<td>29</td>
<td>35</td>
</tr>
<tr>
<td>Age of mother at child’s birth</td>
<td>33</td>
<td>19</td>
<td>34</td>
</tr>
<tr>
<td>Consanguineous marriage</td>
<td>No</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Prenatal damage</td>
<td>preeclampsia</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>Postnatal damage</td>
<td>None</td>
<td>None</td>
<td>Birth asphyxia</td>
</tr>
<tr>
<td>IQ</td>
<td>25</td>
<td>30</td>
<td>28</td>
</tr>
<tr>
<td>CARS Score and Severity</td>
<td>40; Severe</td>
<td>44; Severe</td>
<td>52; Severe</td>
</tr>
<tr>
<td>Comorbidity</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
</tbody>
</table>

### Table 3 Showing pathogenicity prediction of a missense variant

<table>
<thead>
<tr>
<th>Variant</th>
<th>PROVEAN</th>
<th>SNP&amp;GO</th>
<th>PolyPhen2</th>
<th>SNAP2</th>
<th>CADD score</th>
</tr>
</thead>
<tbody>
<tr>
<td>p.N163K</td>
<td>Neutral</td>
<td>Neutral</td>
<td>Benign</td>
<td>Neutral</td>
<td>19.34</td>
</tr>
<tr>
<td></td>
<td>Score: –0.992</td>
<td>Score: 0.381</td>
<td>Score: 0.023</td>
<td>Score: –79</td>
<td>[Raw score 2.010689]</td>
</tr>
</tbody>
</table>
of NLGN4Y gene involved in neurodevelopmental disorder and synaptic functions have been associated with autism.\textsuperscript{17,18} Peripheral blood NLGN4Y gene expression showed an increased risk of autism in children with XYY symptoms.\textsuperscript{7}

In the present study, for the first time in Indian autistic population, we analyzed a cohort of 85 male autistic children under the age of 18 years after screening large neurological disorders population. Our study population records 7.4% of autism over different neurological disorders. Sequencing analysis of the entire exonic region revealed 9 mutations. One is novel missense mutation and remaining 8 (88.9%) are synonymous variants. Out of eight synonymous variants, five (62.5%) were already reported in in-house human SNP databases. Insilco functional effect prediction of a novel missense mutation, p.N163K by PROVEAN, POLYPHEN2, SNAP2, SNP&GO, and CADD shows nonpathogenic effects on the functions of neuroligin 4Y protein. But structural analysis of p.N163K mutant protein shows slight destabilization in the local configuration by positioning on different secondary structure, a turn and it might cause bumps in protein structure due to differences in the size of amino acid. Meanwhile, the mutation introduces a charge; this can cause repulsion of ligands or other residues with the same charge.

Structural and functional prediction of novel missense mutation indicates slight changes related only to the structure of neuroligin 4Y protein, and not to the functions of the protein. Absence of functional analysis of gene was the important limitation of our study.

\section*{Conclusion}

Mutation in NLGN4Y may be an uncommon cause of autism in Indian autistic population. But further detailed functional investigation of neuroligin 4Y gene in autism is important to understand the male predominance of autism and increased rate of autism in males in India autistic population.

\section*{Ethical Approval}
Ethical approval for the study was taken from Institutional Ethical Committee of Shri B.M. Patil Medical College, Hospital and Research Centre, BLDE (Deemed to be University), Vijayapura, [Ref No: BLDE (DU) IEC/337–2018–19]. Informed written consent was obtained from parents/guardians before the collection of blood samples.

\section*{Funding}
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\section*{Conflict of Interest}
None declared.

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