Rett Syndrome without MECP2 Mutation in a Pakistani Girl
Rubina Dad, Humaira Aziz Sawal, Arsalan Ahmad, Muhammad Ikram Ullah, Muhammad Jawad Hassan

ABSTRACT

Rett syndrome is a rare inherited neurodegenerative disease which mostly affects females but has a lethal impact on males. Rett syndrome is mostly caused by mutations of Methyl CpG binding protein-2 (MECP2) gene located on chromosome Xq28.

A 7-year girl from a consanguineous Pakistani family presented with history of abnormal social behavior, tonic colonic seizures, limb'sataxia, intellectual disability, growth retardation and speech abnormalities. Physical and neurological examinations established likely clinical features of Rett syndrome with abnormal electroencephalogram (EEG). Genetic testing of MECP2 gene did not identify any functional nucleotide variation indicating the involvement of another gene mutation in this patient.

A consanguineous case of Rett syndrome did not carry the mutation of MECP2 gene. Due to heterogeneity of the phenotype, it is proposed that there might be involvement of another locus for this disease. In future, targeted next generation sequence can be helpful to identify the causative mutation in this patient.

Key Words: MECP2, Pakistan, Rett Syndrome, Seizures.

Introduction

Rett syndrome (RTT; OMIM 312750) is a rare neurodevelopmental disorder with prevalence of 1 in 15,000 in females. Clinical features include ataxia, loss of speech abilities, seizures onset, intellectual disability, severe developmental delay, cognitive impairments, breathing and swallowing difficulties, chewing and teeth grinding issues and sleep disorders. Methyl-CpG-binding protein 2 (MECP2) gene is responsible for typical RTT in 95% patients and with different RTT feature in 73.2% patients. Other genes like forhead box protein G1 (FOXG1), GRIN1 and KIF1A and cyclin-dependent kinase-like 5 (CDK5) have been demonstrated in congenital RTT and as pathogenic genes of early seizures. The most common causative gene for Rett syndrome is MECP2 (NM_001110792) which is located at candidate region on chromosome Xq28; however, recently some other genes have been identified for this phenotype. A number of mutations in MECP2 have been reported in different ethnic groups and populations. According to various reports, 70%-80% cases of Rett syndrome show mutations in MECP2 gene while other cases are associated with other genes. Depending on the cell type and development phases in the brain, MECP2 imparts variable effects in these processes. RTT is reflected to the failure of functions at different levels, like gene regulation and expression, synaptic function and neuronal circuitry, and during developmental stages.

In the present study, we ascertained a consanguineous family with one affected girl who showed classical features of Rett syndrome and mutation screening of MECP2 gene did not reveal any nucleotide variation. Thus establishing the basis of heterogeneity of this disorder.

Case Presentation

A 7-year old girl visited the hospital, with a history of hyperactive behavior, delayed milestones,
generalized tonic clonic seizures, (onset at 6-month age). Her family history showed a consanguineous relationship between her parents (Fig 1). She was treated previously with medication and remained asymptomatic, followed by a period of relapse. She had remained stable for 2-3 years. Since past three to four years, she had become aggressive.

On recent clinical examination, she was mentally slow with partial loss of fluency of the language and speech. She also developed stereotypic hand movements bilaterally. She had disturbance in breathing when awake and showed impaired sleep pattern. She had growth retardation and with small hands and feet. She was having inappropriate screaming spells, with intense eye communication. Neurological examination revealed normal cranial nerve functions. Manual muscular testing showed weakness of limb and abnormal muscle tone. Her deep tendon reflexes were normal and symmetrically preserved. Her planter response was equivocal on right side and flexor on left side. Electroencephalogram (EEG) showed frequent generalized spikes and wave discharges with mild background slopping and CT scan of head with chest showed bi-frontal atrophy (Fig 2 a, b, c). Urine for mucopolysaccharidosis was negative and Benedict's test for urine glucose was negative. These tests were performed to rule out mucopolysaccharidosis, as there is symptoms overlap. She was given carbamazepine 100mg twice a day with calcium supplements. Genetic testing of MECP2 gene did not identify any functional nucleotide variation associated with this phenotype.

**Table 1: General clinical features of Rett syndrome and comparison with features diagnosed in the patient**

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<thead>
<tr>
<th>Age of onset for Rett syndrome Present study</th>
<th>6-18 months</th>
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<tr>
<td>Present study</td>
<td>6 months</td>
</tr>
<tr>
<td>Primary characteristics Loss of Speech</td>
<td>Yes</td>
</tr>
<tr>
<td>Reduced Hand Growth</td>
<td>Yes</td>
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<tr>
<td>Stereotypic Hand Movements</td>
<td>Yes</td>
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<tr>
<td>Motor Dysfunction</td>
<td>Yes</td>
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<tr>
<td>Autism-Like Features</td>
<td>Yes</td>
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<table>
<thead>
<tr>
<th>Present study</th>
<th>N/A</th>
<th>N/A</th>
<th>N/A</th>
<th>Yes</th>
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<tbody>
<tr>
<td>Peripheral phenotypes Spinal Deformity</td>
<td>Yes</td>
<td></td>
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<td>Principally Scoliosis and Excessive Kyphosis</td>
<td>Yes</td>
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<td>Reduced Bone Mass</td>
<td>Yes</td>
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**Fig 1:** Pedigree representation of family. Filled circle shows affected female and clear squares and circles represent normal individuals. Horizontal line between two individuals demonstrates the relation while vertical lines describe the generations

**Fig 2:** Electroencephalogram and CT scan of affected girl a: EEG shows generalized spike b and c: CT scan of brain showed bilateral atrophic changes

**Discussion**

Rett syndrome is a rare neurodevelopmental disease of childhood. It presents with diverse clinical features like abnormal social interactions, seizures, ataxia, microcephaly, speech and swallowing abnormalities, intellectual disability, delay in growth
and loss of motor movements at later stage of disease. Mutations may be present in MECP2 or other related genes including CDKL5, FOXG1, GRIN1 and KIF1A. In the present case, we report a girl who showed hyperactive social behavior, fits and seizure, loss of speech abilities, limb weakness, intellectual disability and growth retardation. Her EEG showed abnormal wave discharge confirming epileptic fits and CT-scan showed atrophic changes. Supportive diagnosis of Rett syndrome included breathing difficulties, abnormal EEG and seizures, spasticity and ataxia, microcephaly, intellectual disability and delay in growth.

Genetic testing in the present case did not find mutation in MECP2 gene although most common mutations reported worldwide are in MECP2. Mutations in genes other than MECP2, including CDKL5, FOXG1, GRIN1 and KIF1A, are also reported in families with Rett like features due to genetic heterogeneity of this disease.

In Pakistan, very few reports with Rett syndrome epilepsy have been described. Although, diverse features make differential diagnosis very complicated, molecular studies established mutations in MECP2 and FOXG1 in our country.

Next generation sequencing is a remarkable tool for identification of causative genes in heterogeneous disorders like Rett syndrome.

**Conclusion**

This is the fourth case of Rett syndrome reported from Pakistan and this case is without MECP2 gene mutations. In future, other known genes will be sequenced to identify the pathogenic variant through next generation sequencing in this girl.

**Acknowledgments**

We are thankful to the family for participating in this research.

**REFERENCES**