A case of Polymicrogyria with Alpha Dystroglycanopathy presented with milder form with intellectual disability and partial seizure

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**Keywords** : Alpha dystroglycanopathy; complex partial seizures; fukutin; polymicrogyria; developmental delay.

**Background** : Congenital muscular dystrophy (CMD) refers to a group of muscular dystrophies that become apparent in early infancy or at birth. Muscular dystrophies are mostly genetic and a degenerative disease primarily affects voluntary muscles.

**Introduction** : Alpha dystroglycanopathies both phenotypically and genetically are heterogeneous group of disorders and a subgroup of these patients has characteristic brain imaging findings. Material and methods: The case vignette shows a four year old girl presented in the OPD with history of throwing tantrums, delayed developmental milestones, irritability and anger outbursts. She had a history of admission in paediatric neurology indoor with complex partial seizures controlled by tab oxcarbazepine. She was born full term of non-consanguineous marriage by LUCS. There was progressive muscular weakness since early infancy with difficulty in sucking and breathing. No developmental regression was noticed.

**Results** : Her development quotient was found to be 46, plasma ammonia and lactate levels were normal, creatinine kinase was high (314 IU/L). MRI of brain revealed polymicrogyria, white matter changes and subcortical cerebellar cysts. The pattern recognition of MR imaging features may serve as a clue to the diagnosis of alpha dystroglycanopathy although definite diagnosis could be obtained only by muscle biopsy and genetic testing.

**Conclusion** : In Japan, Fukuyama disease is fairly common, second to Duchenne muscular dystrophy but milder form lie this case is rare. The mutation in FKN gene which gives instructions for making a protein called fukutin, which chemically modify a protein alpha-dystroglycan. High index of suspicion and early diagnosis is required to initiate prompt therapy which is mainly supportive with rigorous physiotherapy, antiepileptic drugs, parental and genetic counseling.
Introduction: Alpha dystroglycanopathies are heterogeneous group of disorders and a subgroup of these patients has characteristic brain imaging findings. The clinical features include global developmental delay, contractures, hypotonia and oculomotor abnormalities in all. Other common findings are consanguinity, seizures, macrocephaly or microcephaly, retinal changes and hypogenitalism in majority of patients. 1,2

Case Vignette: A four year old girl presented in the OPD with history of delayed developmental milestones, irritability, history of throwing tantrums and anger outbursts with decreased appetite. She was admitted in paediatric neurology IPD with complex partial seizure with secondary generalization. She was born full term of non-consanguineous marriage by LUCS. It was a valuable pregnancy following primary infertility, six years after marriage. She cried just after birth and there was no significant perinatal insult. There was no regression of developmental milestones. The family history is negative up to three generations. The clinical features included squint, microcephaly (Head circumference below 2 Standard deviations as per age and ethnicity standards) with delayed developmental milestones and complex partial seizure.

CLINICAL AND INVESTIGATION FINDINGS:
On examination, she was found afebrile. Her clinical findings and investigation findings are summarized in Table 1 and Table 2.

Table 1: Clinical features and systemic examination findings of the patient

<table>
<thead>
<tr>
<th>Temperature</th>
<th>98.2 degree F, afebrile</th>
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<tbody>
<tr>
<td>CVS</td>
<td>S1, S2 heard, no murmurs, all peripheral pulses were equally felt without any bruit.</td>
</tr>
<tr>
<td>Respiratory system</td>
<td>Vesicular breath sounds were present, no added sounds.</td>
</tr>
<tr>
<td>Abdominal examination</td>
<td>No Hepatospleenomegaly</td>
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<tr>
<th>CNS</th>
<th>She was alert, conscious and active.</th>
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<tr>
<td></td>
<td>Pupils found 3 mm in diameter and reacting.</td>
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<td></td>
<td>Mild convergent squint.</td>
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<td></td>
<td>Fundus was found to be normal.</td>
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<tr>
<td></td>
<td>No gross abnormalities were found on motor system examination. There was no atrophy, hypertrophy but there was mild spasticity in all four limbs.</td>
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<td></td>
<td>Power was grossly normal.</td>
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<td></td>
<td>Deep tendon reflexes were brisk in both upper and lower limbs.</td>
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<td></td>
<td>Plantar was bilaterally flexor.</td>
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<td></td>
<td>Sensory system examination doesn’t reveal any abnormalities.</td>
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<tr>
<td></td>
<td>Extrapyramidal system- no abnormality was detected.</td>
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<td></td>
<td>Cerebellar signs- absent.</td>
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| Ophthalmological findings | Good visual tracking.  
|                           | Normal anterior segment.  
|                           | Normal fundus.  
| Dermatological findings   | Nothing significant which rules out Junctional epidermolysis bullosa.  
| DQ (Development quotient) | 46  

**Table 2 : The investigation findings of the patient**

| Routine Complete haemogram | Hb 11.3 gm/dl. TLC=477. N55,bE 02, B 00,L 42, M01, ESR=10. PCV=32.0, MCV=88.0, MCH=31.0, MCHC=35.0. RBC=3.6 million, Platelet count=2.2. lakhs. PBS=No abnormal cells has been noted.  
| Plasma ammonia             | 40 (10-65 micromol/L)  
| Plasma lactate            | 16.3 (4.5 to 19.8 mg/d) rules out MELAS (although normal lactic acid doesn’t rule out MELAS unless there is very high CNS involvement.  
| Urine for abnormal metabolites | Not found (rules out storage and urea cycle disorders).  
| Serum creatinine kinase   | 314 IU/L  
| EEG                       | Non specific slowing of waves was found.  
| MRI brain                 | Bilateral fronto-parietal polymicrogyria and tempero-occipital pachygyria with white matter changes has been noted.  
|                           | Cystic changes in cerebellum predominantly in the vermis suggestive of Alpha-Dystroglycanopathy.  
| Echocardiography          | Normal valves and chambers.  
|                           | Normal LV function.  
|                           | No obvious shunt lesion  

**Clinical diagnosis**

Early infantile hypotonia with progressive weakness.
Contractures of hip, knees, interphalangeal joints.
Myopathic facial appearance

High serum CK levels
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**Discussion**: The differential diagnoses made at this stage were (1) Congenital Myopathies, (2) Dystrophinopathies, (3) Emery-Dreifuss Muscular Dystrophy, (4) Metabolic Myopathies, (5) Limb-Girdle Muscular Dystrophy, (6) Spinal Muscular Atrophy, (7) Charcot–Marie–Tooth disease (CMT), (8) Congenital myasthenia etc. But the characteristic MRI findings and raised CPK levels favour the diagnosis of alpha dystroglycanopathy, the Fukuyama congenital muscular dystrophy (milder variant); itself rare, milder form is more uncommon. In major variant the CK level can raise upto 2000-4000 IU/L, here only 314 IU/L. However in Japan, Fukuyama disease is fairly common occurring in approximately 50% of cases. 5,6,7

<table>
<thead>
<tr>
<th>Gene mutation</th>
<th>FKN</th>
<th>LGMD 2M</th>
<th>CMD</th>
<th>CMD-MR</th>
<th>FCMD</th>
<th>WWS</th>
</tr>
</thead>
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<tr>
<td>POMGnT1</td>
<td>LGMD 2O</td>
<td>CMD-MR</td>
<td></td>
<td></td>
<td>MEB</td>
<td>WWS</td>
</tr>
<tr>
<td>POMT1</td>
<td>LGMD 2K</td>
<td>CMD-MR</td>
<td>FCMD</td>
<td>MEB</td>
<td>WWS</td>
<td></td>
</tr>
<tr>
<td>POMT2</td>
<td>LGMD 2N</td>
<td>CMD-MR</td>
<td>FCMD</td>
<td>MEB</td>
<td>WWS</td>
<td></td>
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<tr>
<td>FKRP</td>
<td>LGMD 2I</td>
<td>MDC 1C</td>
<td>CMD-MR</td>
<td>FCMD</td>
<td>MEB</td>
<td>WWS</td>
</tr>
<tr>
<td>LARGE</td>
<td>MDC 1D</td>
<td></td>
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<td>WWS</td>
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Fig. 4: Genetic mutation and clinical phenotypes of alpha-dystroglycanopathies. 8,9,10

The individuals with Ullrich congenital muscular dystrophy has rigid spine with muscular dystrophy (deficiency of selenoprotein N), and integrin-a7 deficiency. The creatine kinase (CK) levels are normal to mildly elevated (≤5 times normal). High CK levels (more than 1000) are found in patients with congenital muscular dystrophy with familial junctional epidermolysis bullosa. CK levels are
mildly to markedly elevated (2-150 times normal) in patients with congenital muscular dystrophy due to abnormal glycosylation or with laminin-α2 mutations (alpha dystroglycanopathy). NCS results are abnormal only in some cases with mutations in laminin-α2, where mild neuropathic changes may be seen. EMG usually shows typical small-amplitude, narrow-duration motor-unit potentials with early recruitment. Prenatal diagnosis had been performed most commonly in families with mutations in laminin-α2 which is expressed in 9-week trophoblasts, in part, because this is the most common congenital muscular dystrophy. Linkage analysis can also be performed but is also at times unreliable, especially in families with partial laminin-α2 deficiency or no brain MRI abnormalities. Muscle biopsy is indicated to confirm the diagnosis and exclude other causes of weakness. In complete laminin-α2 deficiency patients may have severe dystrophic pathology with muscle-fiber degeneration and regeneration, fiber necrosis, and endomyxial and perimysial fibrosis. Immunohistochemical studies show complete loss of staining for laminin-α2.11,12,13

Table 3: Differential diagnosis of the patient.14-17

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<tr>
<td>Most common form (40%) of all cases. May have severe hypotonia, weakness, feeding difficulties, mild intellectual disability and seizure in 30% cases. Periventricular hypomyelination</td>
<td></td>
</tr>
<tr>
<td>3. Integrin-alpha7 deficiency Very rare, presents with hypotonia in infancy with delayed motor milestones. One patient has been followed longitudinally and required noninvasive ventilation at age 8 years and became wheelchair bound at age 12 years.</td>
<td>4. Congenital muscular dystrophy with familial junctional epidermolysis bullosa.[2] Can become severe with blistering lesions and can cause death. Proximal muscle weakness (wheelchair bound) and myesthenic symptoms has been described and may respond to pyridostigmine.</td>
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<tr>
<td>5. Congenital muscular dystrophy with rigid spine (RSMD1). Variable degrees of proximal muscle weakness with hypotonia have been noticed. Carries better prognosis, except few, most patients can eventually walk. In contrast to Ullrich congenital muscular dystrophy, contractures are not present at birth, but they usually develop at age 3-10 years. The most characteristic pattern is spinal rigidity and scoliosis. Contractures of the face, proximal limbs, and finger extensors may also be present. Respiratory insufficiency is common and progressive and may be more severe than limb weakness. Muscle weakness is slowly progressive, no cardio-vascular or MRI abnormalities. Intelligence found normal.</td>
<td>6. Fukuyama congenital muscular dystrophy (mutation in fukutin). Poor sucking, weakness, lack of head control are commonly found. Most patients can walk, progressive weakness and respiratory distress occurs. Cardiac involvement with cardiomyopathy and CCF appears after 10 years. Mild ocular abnormalities, with poor pursuits and strabismus. Can cause retinal detachment, cataracts, severe myopia. Crebellar cysts seen and seizure occur about 50% patients.</td>
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7. LMNA-deficient CMD.
Laminin mutation, mostly present with CMD (Congenital muscular dystrophy) with rigid spine.

8. Muscle-eye-brain (MEB) disease.
Occurs due to genetic mutations. Eye abnormalities and genetic changes are similar to Fukuyama’s disease.

MANAGEMENT

No specific treatment is available for any of the congenital muscular dystrophies. Aggressive supportive care is essential to preserve muscle activity, to allow for maximal functional ability, and to prolong the patient’s life expectancy. The primary neuromuscular concerns include prevention and correction of skeletal abnormalities, such as scoliosis, foot deformities, and contractures, to maintain ambulation. Aggressive use of passive stretching, bracing, and orthopedic procedures, such as spinal fusion, allows the patient to remain independent for as long as possible. Pulmonary complications are the other main concern. Early monitoring and intervention to treat respiratory insufficiency is important because effective therapies can help to improve function and prolong life expectancy. Such therapies include noninvasive bilevel positive airway pressure and/or continuous positive airway pressure or permanent ventilation via a tracheostomy. Cardiac complications are especially common in patients with a mutation in FKRP and occasionally in patients with laminin-α2 deficiency. Treatment of dilated cardiomyopathy with ACE inhibitors and beta-blockers may be necessary. Children with congenital muscular dystrophy may have other neurologic treatment issues, including seizure management, need for supplementary gastric tube feedings, ophthalmologic care, and general medical concerns that occur in profoundly retarded children. As with other hereditary myopathies, a team approach, including a neurologist, pulmonologist, ophthalmologist, cardiologist, orthopedic surgeon, physical medicine specialist, orthotist, and counselors, is required to ensure the best possible care. In patients with CMD with familial junctional epidermolysis bullosa besides the above standard measures, management must include supportive care to protect the skin from blistering, appropriate dressings, and prevention of secondary infections. Activities should minimize skin trauma. Orthopedic surgery is often necessary in patients who live several years with their disease to prevent contractures and scoliosis. According to evidence-based guidelines from the American Academy of Neurology, multidisciplinary care by experienced teams is important for diagnosing and promoting the health of children with CMD.

Patients with alpha-dystroglycanopathies may require prolonged hospitalization. For example, neonates or infants may have progressive disease and have feeding difficulties, cardiopulmonary complications, seizures, or profound mental retardation. Consultation with at least (however not limited to) with the following may prove helpful which (1) Ophthalmologist, (2) Pulmonologist, (3) Cardiologist, (4) Orthopedic surgeon, (5) Epileptologist, (6) Physical and rehabilitation medicine specialist and (7) Dermatologist (patients with CMD with familial junctional epidermolysis bullosa). Older children may need admission for orthopedic care or cardiopulmonary complications. (a) feeding difficulties, (b) respiratory failure, (c) seizures, (d) contractures and/or scoliosis and (e) Blindness. severe disease, such as Walker-Warburg syndrome, patients usually die within the first few years of life. In congenital muscular dystrophy with laminin-α2 deficiency and in some cases of mutations in FKRP, patients occasionally have a relatively normal life span.

CONCLUSION

Most FCMD patients are intellectually disabled and the level is moderate to severe, with IQs ranging from 30 to 50. In a recent study, 62% of patients developed seizures. Among them, 71% had only
febrile seizures, 6% had afebrile seizures from the onset, and 22% developed afebrile seizures following febrile seizures. Most patients had seizures that were controllable with just 1 type of antiepileptic drug, but 18% had intractable seizures that must be treated with 3 medications. The prognosis depends on the type of congenital muscular dystrophy. The Genetic counseling is often helpful to patients and their families to assist in family planning.21

REFERENCES