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A CASE REPORT OF A 2.5-YEAR-OLD GIRL WITH ANGELMAN SYNDROME (AS)

Abstract
Objective
Angelman Syndrome (AS) is a genetically determined syndrome that has a unique behavioral phenotype. This syndrome is described as jerky ataxia and an unusual happy facial expression with pathological laughter. Severe mental retardation is a unique feature of the syndrome, together with microbrachycephaly and abnormal electroencephalographic findings with or without clinical seizures. The patients cannot speak or at most, they have a vocabulary consisting only of a few words. The genetic abnormality of AS has been located on chromosome 15q11-q13. Patients with AS mostly have deletions on the maternally derived allele (75–80%) while some of them show paternal uniparental disomy (~2%) or a rare imprinting mutation developmental disorder caused by deletion of the maternally-inherited chromosome 15q11–13.

A 2.5-year-old girl is presented. Clinical suspicion of AS was raised at the age of 27 months when she presented with mental retardation and epilepsy, absence of speech, inability to gait and paroxysmal episodes of laughter. Moreover, she had facial dysmorphic features such as microbrachycephaly, mid-facial hypoplasia, macrostomia and a prominent mandible. Chromosomal analysis revealed 46 xx with the deletion of 15q chromosome (15q11q13-snrpn/ic) Our patient met the classical phenotype and genotype of AS.

Keywords: Angelman syndrome, microbrachycephaly, Happy facial Phenotype

Introduction
In 1965, Harry Angelman, an English pediatrician, reported 3 children with a similar pattern of mental retardation, seizures, ataxia, easily provoked laughter, absent speech and dysmorphic facial features (1). He called them “puppet children” because of the superficial resemblance to puppets in view of their flat head, jerky movements, protruding tongue and bouts of laughter.

His observations were confirmed over the next decade by other clinicians, and a pejorative term called “happy puppet syndrome” was introduced, not only referring to their gait but also to their joyous facial expression and fits of inappropriate laughter(2). Williams and Frias proposed the term “Angelman’s syndrome”.

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Over 450 AS individuals have now been reported including familial cases. The prevalence of AS is estimated to be around 1/10,000–1/20,000 (3,4). AS is a clinical diagnosis that can be confirmed by either cytogenetic or DNA testing in about 80–85% of the cases. Patients with AS have a typical phenotype. Features of AS frequently become apparent at 1–4 years of age, the average age at diagnosis is 6 years and the syndrome is characterized by dysmorphic cranio-facial features (micro-brachycephaly, mid-facial hypoplasia, deep set eyes, macrostomia and prominent mandible), jerky ataxia and an unusual happy facial expression with pathological laughter and Severe mental retardation with or without clinical seizures (5-9).

We report a patient with Angelman syndrome and describe its different features to help our physicians to have a more vivid picture of this condition.

Case report
The patient, a 2.5-year-old girl, was the first child of healthy first-cousin parents. She was born at term gestation by spontaneous vaginal delivery and weighed normal for gestational age. She had a positive family history of epilepsy and mental retardation in her aunt and her cousin.

Facial features included microcephaly (head circumference: 45.5 cm, below the 3rd percentile) with a wide mouth and protruded tongue, as well as mandibular prognathism. She had a happy disposition (already noted in infancy) with paroxysms of laughter (figure 1).

Generalized tonic clonic seizures were presented from the age of 7 days which were treated promptly with phenobarbital. Seizures continued until the anticonvulsants were changed to primidon and clobazam. Concomitant EEG findings were markedly abnormal with generalized spike and poly spike waves. At the time of the diagnosis (age of 27 months), the patient demonstrated a marked neuro-developmental delay with severely impaired communication and lack of any recognizable speech. She was able to sit, but could not walk independently. A performed CT scan at that age was normal. Chromosomal analyses with karyotype (GTG) and fluorescent in situ hybridization (FISH) techniques revealed 46 xx with deletion of 15q chromosome (15q11q13-snrpn/ic) (Figure 2-3).

Discussion
Our patient in this study was diagnosed at the age of 27 months with facial features including microcephaly and mental retardation. Other symptoms were absence of speech, inability to gait and paroxysmal episodes of laughter with a wide mouth, a protruded tongue and mandibular prognathism. She had a happy disposition with paroxysms of laughter (10-12).

In other studies, the features of AS frequently become apparent at 1–4 years of age with the average age at diagnosis being 6 years (10-12). The facial features evolve over the first 5 years of life. Typical features include brachycephaly and a head circumference below the 25th percentile (13). A horizontal occipital groove is found in 35% of the patients. The mouth is large, giving rise to widely spaced but otherwise normal teeth.

The chin is pointed and there is mandibular prognathism. The eyes are deep set and often blue. Strabismus is seen in about 40% of the patients (13). Severe visual problems are not common. Coarsening of facial features with increasing age has been reported, with marked mandibular prognathism, a pointed chin, macrostomia and a pronounced lower lip (14-16).

All patients have severe mental retardation and delayed motor milestones. Jerky movements become apparent during the first few months and motor delay is obvious by nine months of age. The jerky movements, tongue thrusting, mouthing, and hand-flapping when walking, are all characteristic. Reflexes are brisk and the gait is slow, ataxic and stiff-legged with the characteristic posture of raised arms with flexed wrists and elbows. Almost all patients are ‘happy’ and smile frequently. Their laughter is provoked by minimal stimuli and is occasionally inappropriate (13-15).

In this patient, seizure was present from the age of 7 days and was treated promptly with phenobarbital. Seizures continued until the anticonvulsant was changed to primidon and clobazam. Concomitant EEG findings were markedly abnormal with spike and
poly spike generalized waves. Studies demonstrate that epileptic seizures occur in about 80% of the patients with AS. Age at onset varies from 1 month to 5 years (mean 2–3 years). The initial symptoms of epilepsy are febrile convulsions in infancy in about 40% of the patients (17-18).

In this patient, EEG findings were markedly abnormal with generalized spike and poly spike waves. There are specific EEG patterns in AS patients which may appear in isolation or in various combinations either in the same EEG recordings or at different times in the same patient (18-20).

Computerized tomography (CT scan) findings in two studies were normal or showed cerebral atrophy and ventricular dilatation in a minority of the patients (13-14). One report described an abnormally convoluted surface area of the cortex in the parietal lobe of the Sylvian fissure in the supramarginal gyrus, using specific techniques in MRI (Magnetic Resonance Imaging)(21).

There is no specific therapy for AS patients. Epileptic seizures often need AEDs (Anti Epileptic Drugs) and good results are achieved with Valproic Acid and Clonazepam. Physiotherapy is very important for AS patients to keep them mobile as long as possible and to minimize orthopedic interventions. The use of non-verbal communication is potentially useful for them, but attempts to train the use of signing to augment their speech may be unsuccessful due to their poor imitation skills and possible motor organizational difficulties (22).

In conclusion, This study demonstrated a typical manifestation of Angelman syndrome with facial features, mental retardation, delayed motor and seizures with EEG findings.

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![Figure 2](image1.png)

**Fig 2.** Chromosome study of the patient on the basis of GTG technique: Twenty metaphase spreads were studied on the basis of GTG technique at 450-500 band resolution, revealing 46 chromosomes with possible deletion in long arm of chromosome 15 at q11q13 band.

![Figure 3](image2.png)

**Fig 3.** Chromosome study of the patient on the basis of fluorescent in situ hybridization: Fluorescent in situ hybridization was performed on metaphase sreads using the cytocell prader-willi/Angelman (SNRPN/IC) probe. Conclusion: 46XX, ish del(15)(q11q13)(SNRPN/IC) Compatible with prader-willi/Angelman syndrome.
References

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