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***SATB2*-associated syndrome caused by a novel *SATB2* mutation in a Chinese boy: A case report and literature review**

Zhu Yy *et al*. A novel *SATB2* mutation for SAS

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**Abstract**

BACKGROUND

Special AT-rich sequence binding protein 2 (SATB2)-associated syndrome (SAS; OMIM 612313) is an autosomal dominant disorder. Alterations in the *SATB2* gene have been identified as causative.

CASE SUMMARY

We report a case of a 13-year-old Chinese boy with lifelong global developmental delay, speech and language delay, and intellectual disabilities. He had short stature and irregular dentition, but no other abnormal clinical findings. A *de novo* heterozygous nonsense point mutation was detected by genetic analysis in exon 6 of *SATB2*, c.687C>A (p.Y229X) (NCBI reference sequence: NM\_001172509.2), and neither of his parents had the mutation. This mutation is the first reported and was evaluated as pathogenic according to the guidelines from the American College of Medical Genetics and Genomics. SAS was diagnosed, and special education performed. Our report of a SAS case in China caused by a *SATB2* mutation expanded the genotype options for the disease. The heterogeneous manifestations can be induced by complicated pathogenic involvements and functions of SATB2 from reviewed literatures: (1) SATB2 haploinsufficiency; (2) the interference of truncated SATB2 protein to wild-type SATB2; and (3) different numerous genes regulated by SATB2 in brain and skeletal development in different developmental stages.

CONCLUSION

Global developmental delays are usually the initial presentations, and the diagnosis was challenging before other presentations occurred. Regular follow-up and genetic analysis can help to diagnose SAS early. Verification for genes affected by *SATB2* mutations for heterogeneous manifestations may help to clarify the possible pathogenesis of SAS in the future.

**Key words:** Special AT-rich sequence binding protein 2; *SATB2*-associated syndrome; Global developmental delay; Developmental speech and language delay; Case report

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**Core tip:** Our findings contribute to a growing list of special AT-rich sequence binding protein 2 (*SATB2*) mutations associated with SATB2-associated syndrome, which is a rare autosomal dominant disorder. The diagnosis was quite challenging when only developmental delays occurred without other manifestations. The heterogeneous manifestations can be induced by complicated pathogenic involvements and functions of SATB2 from reviewed literatures: (1) SATB2 haploinsufficiency; (2) the interference of truncated SATB2 protein to wild-type SATB2; and (3) different numerous genes regulated by SATB2 in brain and skeletal development in different developmental stages. We think our case can help the practitioners to learn more about SATB2-associated syndrome.

**INTRODUCTION**

Special AT-rich sequence binding protein 2 (SATB2)-associated syndrome (SAS; OMIM 61231) is a multisystem disorder characterized by neurodevelopmental delays, behavioral issues, palatal and dental anomalies, craniofacial abnormalities, and frequent skeletal pathology[1]. It was first reported in a 16-year-old male in 1989 and termed Glass Syndrome at the time[2].The frequency of SAS has been estimated in individuals with undiagnosed intellectual disability (ID)/developmental delay (DD) to be 0.24%-0.3%[3,4].The diagnosis of SAS is likely to be inadequate and should be considered in children with ID, speech delay, cleft or high-arched palate, and abnormal dentition[5]. Up to July 2020, 194 individuals with SAS from more than 15 different countries had enrolled at a website for SAS (www.SATB2gene.com). The acronym has also been used to describe the signs: Severe speech anomalies, Abnormalities of the palate, Teeth anomalies, Behavioral issues with or without bone or brain magnetic resonance imaging (MRI) anomalies, and age of onset before 2 years of age (S.A.T.B.2.)[1].

For representative cases in a typical stage with the classic signs, the possible diagnosis is easy to consider. No specific laboratory tests are available for SAS, and non-specific elevation of alkaline phosphatase levels has been reported in some individuals with osteopenia[4,6].Thus, early diagnosis is quite challenging in infants with only DD observed. The mean age at diagnosis has been reported to be 6.6 years (range: 1 wk to 34 years)[7].

Alterations in *SATB2* have been identified as pathogenic causes, including contiguous deletions, intragenic deletions and duplications, translocations with secondary gene disruption, missense point mutations, and gonadal/somatic mosaicism[1,8-11]. SATB2, which is a kind of matrix attachment region–binding transcription factor and has a homeodomain, a CUTL domain, and two CUT domains, plays a key role in brain and skeletal development[12,13]. Human patients with SATB2 haploinsufficiency have shown some neurological symptoms, such as cognitive deficits, developmental delay, and developmental speech and language problem[14]. Here, we report a case of SAS confirmed by a novel nonsense mutation in *SATB2* that is first reported here.

**CASE PRESENTATION**

***Chief complaints***

The proband (Figure 1A) was a 13-year-old Chinese boy with lifelong global DD. Brain development evaluations and genetic analysis were planned for the current medical visit after referring to local medical geneticists because the family hoped to have one more child.

***History of present illness***

There was no exposure to medications or alcohol during the pregnancy, and he was born spontaneously at 35 wk and in an incubator for 3 wk. He presented with delayed developmental milestones: unassisted sitting at about 1 year, unassisted standing at about 20 mo, and unassisted walking at about 2 years. He could speak only one word at 2 years old. He was admitted to the local hospital at that time, and brain MRI showed abnormal long T2 signals in the white matter, but electroencephalography was normal. He was diagnosed with cerebral palsy and brain dysplasia and did not receive any rehabilitation training before receiving special education from 6-years-old. He can eat and go to the toilet by himself but still speaks monosyllabic words and can hardly communicate with others. He experiences less sleep and is excitable.

***History of past illness***

He showed no other health problems in the past.

***Personal and family history***

He is the only child of nonconsanguineous parents without marked family histories.

***Physical examination***

Upon physical examination, the patient weighed 28.3 kg (< 3rd percentile), his height was 140.5 cm (< 3rd percentile), and the occipitofrontal circumference was 50 cm (normal range). He was conscious and had a normal gait. He also presented irregular dentition but no special facial features. Patellar reflex was normal, and Babinski signs were negative. The hand-alternating movement test, heel-knee-tibia test, and intelligence quotient value could not be completed because of non-cooperation with the instructions.

***Laboratory examinations***

Blood tests were normal, including ceruloplasmin, lactic acid, ammonia, and liver and kidney functions. Screening for genetic and metabolic diseases in the blood and urine was not suggestive of disease.

The next generation sequencing-based trio whole exome sequencing plus copy number variation sequencing was performed for genetic analysis (the process was performed by Beijing Chigene Translational Medicine Research Center). The EDTA-treated peripheral blood were collected with informed consent of the patient’s guardians. The DNA was extracted using the Blood Genome Column Medium Extraction Kit (Kangweishiji, China) according to the manufacture instructions and was subjected to quality control using Qubit 2.0 fluorimeter and electrophoresis with 0.8% agarose gel for further protocol. The whole exome library was constructed using xGen Exome Research Panel v1.0 (IDT, Iowa, United States) that consists of 429826 individually synthesized and quality-controlled probes, which targets 39 Mb protein-coding region (19396 genes) of the human genome and covers 51 Mb of end-to-end tiled probe space. High-throughput sequencing was performed by Illumina NovaSeq 6000 series sequencer (PE150). A *de novo* heterozygous nonsense point mutation in exon 6 in *SATB2,* c.687C>A (p.Y229X) (NCBI reference sequence: NM\_001172509.2) was detected in the proband. Direct sequencing validated the detected missense mutations. PCR was performed to amplify the exon with the detected mutation and its neighboring introns using the forward primer: 5’-TAGTTTTAAGGGAGCCAACTAGGA-3’ and the reverse primer: 5’-TCAGTTTACAGGAGAGTGACTGTTT-3’. PCR products were purified using micro concentrating centrifugal filter columns (Millipore, Bedford, MA, United States), sequenced by ABI 3730XL (Applied Biosystems, Forster City, CA, United States), and analyzed by DNASTAR software.

The proband had the mutation that neither of his parents had (Figure 1B). The detected heterozygous mutation c.687C>A (p.Y229X) is near the end of coding region for the CUTL domain and leads to a stop codon. We think the mutation c.687C>A (p.Y229X) leads to an abnormal transcription and then a truncated protein, like a nearby mutation c.715C>T (p.R239X) that had been reported to lead to an abnormal transcription and a truncated protein[3,15,16]. We did not investigate the mutation in control groups, but the minor allele frequency for the mutation was not found in the dbSNP database (https://www.ncbi.nlm.nih.gov/snp/), 1000 Genomes Project[17],Virtual Chinese Genome Database[18], or Genome Variation Map[19]. The effect of the mutation was evaluated according to the guidelines of the American College of Medical Genetics and Genomics[20], and the mutation was considered pathogenic (Table 1). The mutation is the first reported for SAS.

***Imaging examinations***

An electroencephalography was normal, but brain MRI revealed multiple small cystic lesions in the white matter near the posterior horns of the right lateral ventricle and long T2 signals in the white matter adjacent to the bilateral posterior horns of the lateral ventricles (Figure 2A).

**FINAL DIAGNOSIS**

SAS was diagnosed.

**TREATMENT**

After diagnosis, special education in local welfare institutions and symptomatic treatment were performed (for example: oral chloral hydrate for sleeping problems if necessary).

**OUTCOME AND FOLLOW-UP**

At 6 mo follow-up, no speech improvements were observed.

**DISCUSSION**

For SAS patients, preterm birth was reported quite rarely[1,2]. The first reported patient was delivered at 32 wk[2]. But *SATB2* is not a candidate gene for spontaneously preterm birth for now. For our case, he was delivered at 35 wk without obvious causes, and we did not find associations between the preterm birth and *SATB2*. Many neurodevelopmental problems are associated with prematurity compared to full term infants, including DD, language problems, and poorer academic performance in the future. Some developmental problems may be corrected if interventions were performed early[21]. Differentiating underlying causes for DD early in premature infants can help to make more accurate treatment maneuvers. SAS includes a constellation of manifestations that do not occur simultaneously or necessarily (Table 2).

***Severe speech anomalies***

All SAS patients have DD/ID, 95% have speech delay[22],82% have a limited vocabulary under ten words or absent speech by 10-years-old, and their first words are spoken at a median of 24 mo of age (range 10-144 mo)[7]. In three patients from Japan, meaningful words had not been acquired at 5-years-old[23]. The process usually did not regress after gaining some abilities. Only one case with an 8.6-Mb deletion of 2q32.2-q33.1, including *SATB2,* was reported to have ID regress from mild to severe and speech from poor to absent between the ages of 6 and 12 years[24]. For our case, DD and limited speech were considered to be caused by the prematurity, and it was hoped that he would develop to normal in the future when he was referred to the hospital in the first few years. Therefore, we think the attention to the prematurity history may be one cause for the delayed diagnosis for the patient.

***Abnormalities of the palate***

Seventy-six percent of individuals with SAS have palatal abnormalities such as cleft palate, high-arched palate, and bifid uvula. In 42% of SAS individuals, micrognathia is found[22]. For our case, no such abnormalities were found.

***Teeth anomalies***

Thirty-six percent of individuals with SAS have abnormal upper central incisors. In addition, 36% have dental crowding, 16% have hypodontia, 6% have delayed primary dentition, and/or 4% have diastema[22].Delayed permanent root formation, delayed or missing second bicuspids, and malformed teeth have been found on radiography[25].Multiple odontomas have been reported as an occasional manifestation in SAS patients[26]. In our case, dental crowding and irregular dentation were found, and he still did not grow any permanent teeth.

***Behavioral issues with or without bone or brain MRI anomalies***

A broad spectrum of behavioral issues has been reported, including pleasant personality, irritability, hyperactivity, sleep problems, sensory issues, stereotyped repetitive movements, and Rett syndrome-like phenotypes[3,7,22,27,28].Nonspecific brain abnormalities detected by MRI are found in half of affected individuals, including ventricle enlargements, thin corpus callosum, enlarged perivascular spaces, and abnormal myelinization in white matter[7,22]. Some Pectus, finger, or spine deformities have been reported in a few affected individuals, some with concurrent osteopenia[8,22]. In our case, the patient slept less, and brain MRI indicated periventricular leukomalacia and poor myelinization, which is relatively common in subjects with a history of preterm or neonatal respiratory distress syndrome, and showed normal corpus callosum without obvious physical abnormalities (Figure 2B).

***Age of onset before 2 years***

Individuals with SAS fail to reach normal developmental milestones from infancy. Some other signs may exist from infancy, such as mild but nonspecific facial dysmorphism, hypotonia, feeding difficulties, clinical seizures, and abnormal muscle strength[22].Growth restriction, severe heart defects, small or undescended testicles, inguinal hernias, hypospadias, and thin skin or hair have been reported in several cases with large deletions including *SATB2*[22]. In our case, the patient had short stature, though genetic analysis did not reveal large gene deletions.

According to the gene database for SAS (https://satb2gene.com/pros-molecular-data/), 194 variants have been identified, and single nucleotide variants causing a premature stop codon were the most frequent type (46%) followed by missense variants (25%), large deletions > 1 Mb (19%), and small deletions (12%). According to the published data, 120 unique variants have been identified in *SATB2*, and the most common ones were single nucleotide variants inducing the occurrence of a stop codon (42.5%), and next were the missense variants (25.8%). About 74.2% of missense variants are located in CUT1, CUT2, or homeodomain DNA protein domains[12]. The mutation detected in our case (c.687C>A (p.Y229X)) is a nonsense point mutation that has not been reported previously and located in the CUTL domain.

The phenotypes are heterogeneous according to the fragment deletion size and locations of mutations in *SATB2* [12]. For example, abnormal myelination and/or white matter abnormalities were relatively common (26%) in patients with pathogenic nonsense, missense, and frameshift variants, while abnormalities in heart, testicles, skin, or hair were reported in patients with large fragment deletions[7,22].In addition, even the same genotypes of p.R239\* that have been detected in different individuals have phenotypes that were not identical (from Genotype-Phenotype database at https://satb2gene.com/pros-molecular-data/).

The reasons for the heterogeneous manifestations for SAS can be the complicated pathogenic involvements and functions of SATB2. SATB2 is a kind of matrix attachment region-binding transcription factor, with high levels of expression in the brain, including the cerebral cortex and spinal cord, and plays a role in central nervous system development[29]. SATB2 haploinsufficiency can be the cause for intellectual disability. Mutant SATB2 protein appears functionally inactive because of the disrupted associations with chromatin or matrix[3,14]. The oligomerization of the N-terminal domain of SATB1, another SATB family protein, is critical for DNA-binding affinity[30]. The truncated protein with left N-terminus and CUTL domain in a patient with a heterozygous nonsense mutation of c.715C>T (p.R239X) has been shown to localize to the nucleus and interferes with the normal activity of wild-type SATB2 protein by forming a dimer with wild-type SATB2[16]. In our case, the mutation is near the reported mutation of c.715C>T (p.R239X), induces a stop codon at the end of the CUTL domain, and can leave an unaffected N-terminus and oligomerization domain of SATB2. This may disturb the DNA binding and gene regulation like the reported mutation of c.715C>T (p.R239X).

SATB2 is involved in transcription regulation and chromatin remodeling. SATB2 can activate *UPF3B* transcription, and defects in *UPF3B* in humans can induce cognitive deficits and craniofacial dysmorphisms[16]. SATB2 regulates a transcriptional network of multiple key determinants for skeletal development by activating or repressing DNA bound protein or enhancing the activity of other DNA binding proteins[13]. Three gene sets related to or regulated by SATB2 have been analyzed at different stages of development; the sets include many genes contributing to schizophrenia and educational attainment[31].Other research has shown that SATB2 interacts with different protein networks at developing and adult stages in the cortex, indicating that the SATB2 function shifts in the cortex over a lifetime[13]. In the mouse, Satb2 regulates regionalization of the retrosplenial cortex by controlling Nr4a2 and Ctip2 during development[32], and loss of Satb2 in cortex and hippocampus contributed to abnormal behavior[33]. Satb2 was involved in the specification of upper-layer neuron specification in the neocortex in mouse by regulating expression of specific genes[34]. In the developing cerebral cortex, Satb2 regulated the formation of corticocortical connections by repressing the expression of *Ctip2*, which is a transcription factor involved in the extension of subcortical projections from cortical neurons[35,36].

The SATB2 function shift at different developmental stages and complex interactions with related proteins may be involved in the pathogenesis of variable manifestations in SAS patients.

Diagnosis for SAS can be quite challenging in infants with only DD, hypotonia, feeding difficulties, and palatal issues. Cerebral palsy may be considered at this time, especially for those who were preterm or had neonatal asphyxia, but nonprogressing motor developmental abnormalities during follow-up are the main characteristic of cerebral palsy[37]. Leukodystrophy is usually the initial diagnosis for those with some neurological manifestations and abnormal findings in white matter on brain MRI and can occur at different ages, from newborn to adult. The initial clinical manifestation of leukodystrophies is often nonspecific, such as motor impairment, cognitive impairment, and seizures. Further diagnosis of leukodystrophies constitutes many kinds of rare heritable disorders and is challenging in most cases[38].Angelman syndrome and related disorders can be considered during infancy, which are characterized by severe cognitive disability, motor dysfunction, speech impairment, and hyperactivity, but inappropriate happy demeanor may help diagnose[20,39]. Dental and behavioral problems over time may indicate SAS during follow-up. KBG syndrome manifests dental abnormalities (macrodontia of upper central incisors), DD, and other characteristic features that help in diagnosis, including distinctive craniofacial features, such as triangular face, broadened nasal bridge, thin upper lip, and synophrys[20,40].

Today, next generation sequencing technologies, especially whole exome sequencing and whole genome sequencing, are available for clinical genetic diagnosis and differential diagnosis of heritable diseases, especially for distinct differential diagnosis[41]. Cerebral palsy and leukodystrophy are clinical diagnoses, but the etiologies vary and are not fully known. Approximately one-third of cases do not have traditional risk factors, such as prematurity, hypoxic–ischemic injury, placental insufficiency, and prenatal infection, and genetic mutations may be involved in the pathogenesis of a substantial proportion of cases of cerebral palsy[37]. Distinct diagnoses of leukodystrophies have significantly increased due to genetic diagnostic tests[38].Genetic testing of *UBE3A* for Angelman syndrome and*ANKRD11* for KBG syndrome can help to differentiate the diseases from SAS[39,40].

No specific therapy or guidelines are available for SAS, and regular evaluations after diagnosis and treatment of symptoms are recommended. Specialists in different fields deal with sleep disturbances, seizures, feeding difficulties, behavioral issues, scoliosis, tibial bowing, and joint contractures, among others. Developmental support/special education is referred for DD/ID[20].Prompts for restructuring oral muscular phonetic targets and aggressive speech therapy (3 ×/wk) at 2-years-old have shown improvements in speech over time, and some communication devices, such as picture exchange communication system, may provide additional help for SAS patients[1].

**CONCLUSION**

We confirmed SAS by the genetic analysis and collected the clinical features. A new variant was found. Because of phenotypic heterogeneity, limited physical findings, and lack of specific laboratory tests, SAS can be difficult to diagnose in infants with DD but no other signs. Surveillance should focus on speech development, dental status, and behavior issues. Next generation sequencing can help differentiate underlying diseases and predict the appropriate prognosis, and we recommend genetic testing for children who show delays over 6 mo compared with normal development milestones in gross movement and/or speech and language development in clinical practice. In future studies, verification for different gene sets that are affected by *SATB2* mutations for heterogeneous manifestations may help to clarify the possible pathogenesis of SAS.

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**Footnotes**

**Informed consent statement:** Consent was obtained from the patient and his/her relatives for the publication of this report and any accompanying images.

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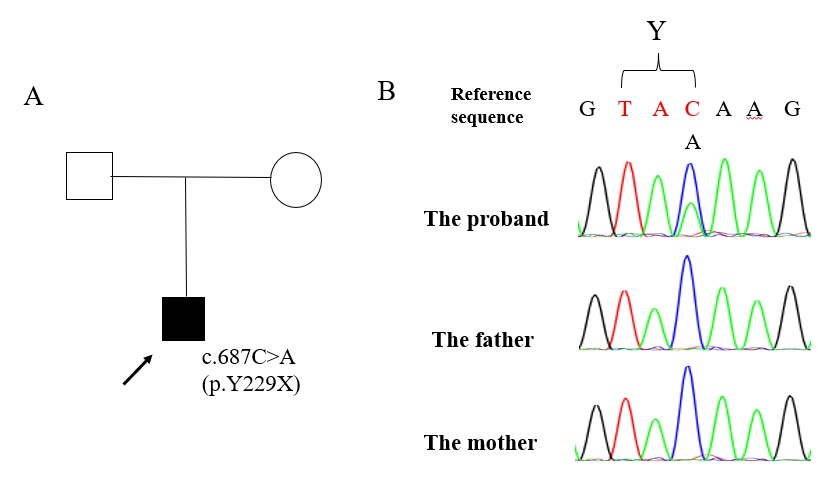
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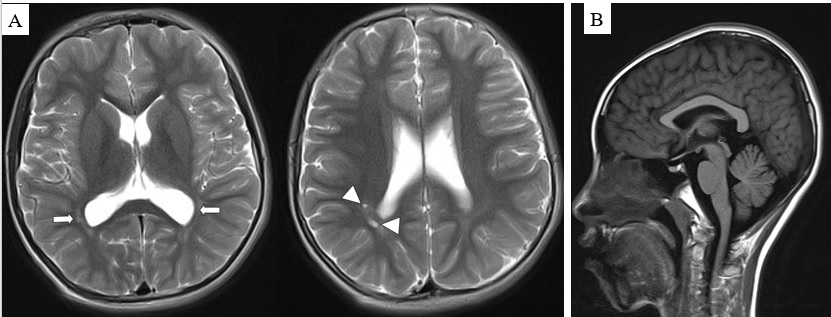
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**Figure Legends**



**Figure 1 The family pedigree and mutations detected in *SATB2*.** A: The pedigree of the family with *SATB2*-associated syndrome (SAS). The arrow indicates the proband; the parents have no signs of *SATB2*-associated syndrome; B: The mutations detected in the family. The mutation is *de novo* in the proband, whereas the parents are wild-type.



**Figure 2 Brain magnetic resonance imaging images.** A: Multiple small cystic lesions in the white matter adjacent to the posterior horns of the right lateral ventricle (arrowheads) and long T2 signals in the white matter adjacent to the bilateral posterior horns of the lateral ventricles were found in transversal T2-weighted images(arrows); B: The corpus callosum was found normal in sagittal fluid-attenuated inversion recovery image.

**Table 1 Functional evaluation of the c.687C>A (p.Y229X) mutation in** special AT-rich sequence binding protein 2 **according to the guidelines from American College of Medical Genetics and Genomics[15]**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Functional evaluation | **Weight for pathopoiesis** | **Pathogenic criterion for the mutation** | | |
| PVS1 (very strong) | Null variant in *SATB2* where loss of function is a known mechanism of the disease | | |
| PM2 (moderate) | Absence in population database | | |
| PP3 (supporting) | Multiple lines of computational evidence support a deleterious effect on the gene/gene production | | |
| PolyPhen-2 | SIFT | MutationTaster |
| Probably damaging | Damaging | Disease causing |
| Final evaluation | Pathogenic | | | |

*SATB2*: Special AT-rich sequence binding protein 2.

**Table 2 Main clinical findings for** special AT-rich sequence binding protein 2**-associated syndrome (according to** **lecture[22] and https://satb2gene.com)**

|  |  |  |
| --- | --- | --- |
| **Signs for SAS** | | **Reported frequency (%)** |
| Demographics | Gender (M:F) | 3:2 |
| Age < 4 yr | 31 |
| Age 4-10 yr | 37 |
| Age 10-18 yr | 19 |
| Adults | 13 |
| Severe speech anomalies | DD/ID | 100 |
| speech delay | 95 |
| Abnormalities of the palate | cleft palate, high-arched palate, and bifid uvula | 76 |
| micrognathia | 42 |
| Teeth anomalies | abnormal upper central incisors | 36 |
| dental crowding | 36 |
| hypodontia | 16 |
| delayed primary dentition | 6 |
| diastema | 4 |
| Behavioral issues with or without bone or brain MRI anomalies | Feeding difficulties | 39 |
| Growth restriction | 34 |
| Enlarged ventricles | 12 |
| Agenesis of corpus callosum | 5 |
| Age of onset before 2 yr | — | — |

SAS: Special AT-rich sequence binding protein 2-associated syndrome; MRI: magnetic resonance imaging; DD/ID: developmental delay/intellectual disability; M: Male; F: Female.