

A Longer Polyalanine Expansion Mutation in the *ARX* Gene Causes Early Infantile Epileptic Encephalopathy with Suppression-Burst Pattern (Ohtahara Syndrome)

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Early infantile epileptic encephalopathy with suppression-burst pattern (EIEE) is one of the most severe and earliest forms of epilepsy, often evolving into West syndrome; however, the pathogenesis of EIEE remains unclear. *ARX* is a crucial gene for the development of interneurons in the fetal brain, and a polyalanine expansion mutation of *ARX* causes mental retardation and seizures, including those of West syndrome, in males. We screened the *ARX* mutation and found a hemizygous, de novo, 33-bp duplication in exon 2, 298_330dupGCGGCA(GCG)_n, in two of three unrelated male patients with EIEE. This mutation is thought to expand the original 16 alanine residues to 27 alanine residues (A110_A111insAAAAAAAAAAAA) in the first polyalanine tract of the *ARX* protein. Although EIEE is mainly associated with brain malformations, *ARX* is the first gene found to be responsible for idiopathic EIEE. Our observation that EIEE had a longer expansion of the polyalanine tract than is seen in West syndrome is consistent with the findings of earlier onset and more-severe phenotypes in EIEE than in West syndrome.

Early infantile epileptic encephalopathy with suppression-burst pattern (EIEE) is one of the most severe and earliest forms of epilepsy and has been known as “Ohtahara syndrome” since Ohtahara et al. first reported it in 1976.¹ Frequent tonic seizures or spasms start mainly in the neonatal period, with a specific finding of electroencephalography (EEG)—namely, suppression-burst pattern characterized by high-voltage bursts alternating with almost flat suppression phases at an approximately regular rate.² Cerebral dysgenesis—including hemimegalencephaly, Aicardi syndrome, and porencephaly—is often accompanied by EIEE; however, the etiology is heterogeneous, and cryptogenic or idiopathic EIEE has been reported only in a small number of patients. Familial occurrence of EIEE has been reported only in a family with Leigh encephalopathy.³ The transition from EIEE to West syndrome, which is characterized by tonic spasms with clustering, arrest of psychomotor development, and hypsarrhythmia on EEG, occurs in 75% of patients with EIEE,⁴ and a common pathological mechanism between them has been suggested. Recently, two novel genes, *ARX* and *CDKL5*, have been found to be responsible for X-linked West syndrome or infantile spasms (ISSX [MIM 308350]).^{5–8}

ARX, the aristaless-related homeobox gene (GenBank accession number NM_139058), is located in the human chromosome Xp22 region and as a transcription factor is essential for the development of cerebral interneurons.⁹ Mutation of *ARX* causes a wide variety of phenotypes, from severe brain malformations like hydranencephaly or

lissencephaly lacking nonpyramidal or GABAergic interneurons to nonsyndromic mental retardation, with a strong correlation between genotype and phenotype.^{10,11} To identify the gene(s) underlying EIEE, we screened the *ARX* mutation in three male patients with EIEE defined by brief tonic seizures and a suppression-burst pattern on EEG with unknown etiology. Female patients were excluded, because even the null mutation of *ARX* causes less severe phenotypes than West syndrome.¹⁰ All patients showed a transition from EIEE to West syndrome and severe developmental delay. Clinical information regarding the patients in whom the *ARX* mutation was found was as follows: patient 1 was the first child of unrelated healthy parents. He was born normally at term without asphyxia. He suffered his first seizure 40 min after birth. His eyes deviated to the left with loss of consciousness, followed by brief clonic movement, predominantly of the left leg and arm. The seizure ceased after ~1 min and recurred four times. Physical examination demonstrated a microcephalus (fig. 1A). Findings from his blood and cerebrospinal fluid examinations were normal. A CT scan of his head also showed normal results. EEG examination demonstrated a suppression-burst pattern that was compatible with EIEE (fig. 1B). His seizure types shifted from (1) eye deviation followed by brief clonic movement to (2) brief tonic seizures and pedaling movements and then to (3) brief tonic spasms in clusters. He tended to cry with an opisthotonic posture. After 1 mo, EEG demonstrated a transition to hypsarrhythmia, suggesting West syndrome,

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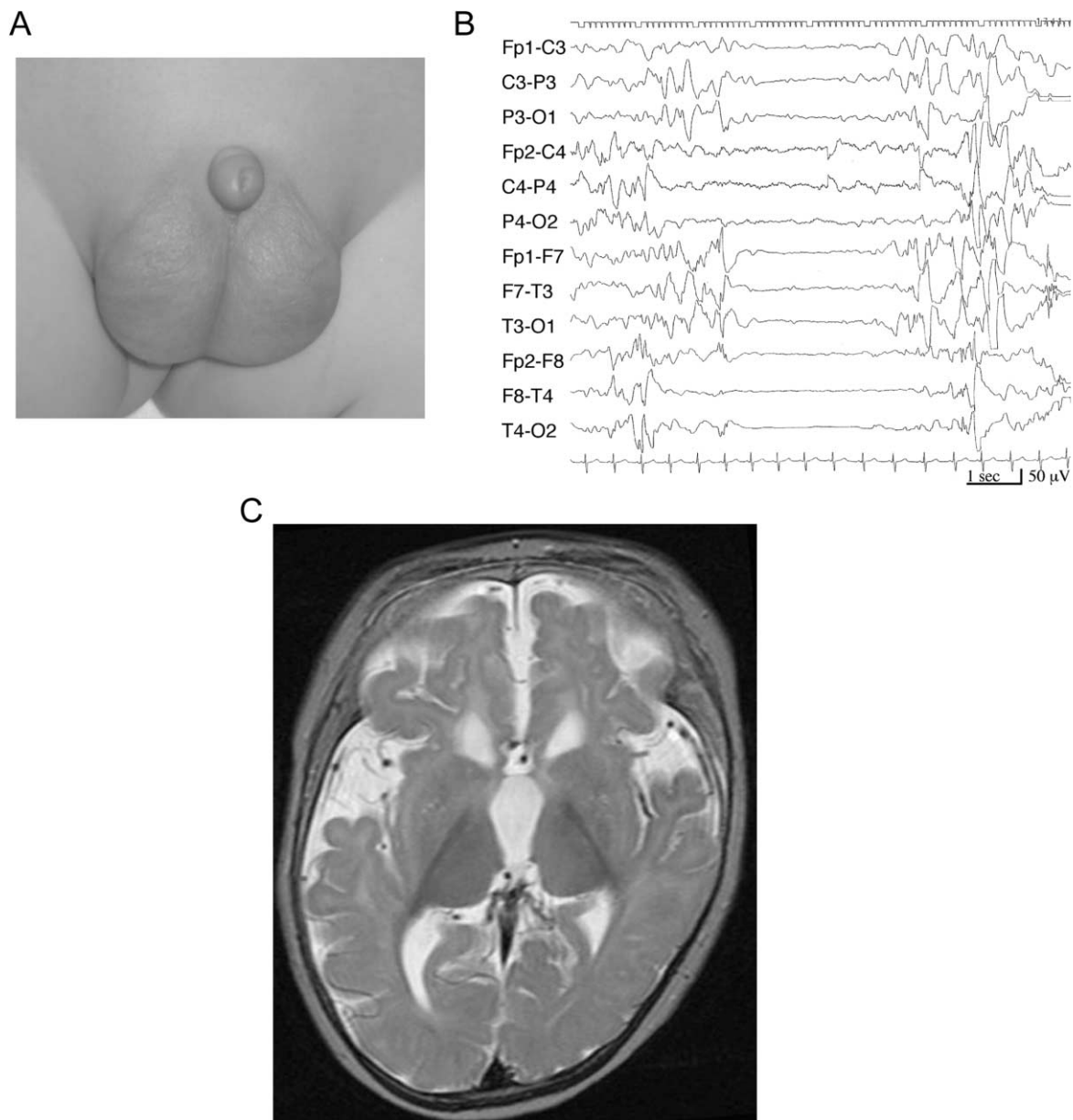


Figure 1. *A*, Photograph of patient 1's external genitalia, showing a small penis with an apparently normal scrotum. *B*, Interictal EEG of patient 1, showing an asynchronous pattern, with high-voltage slow wave bursts mixed with spikes and sharp waves, and a low-voltage suppression background. *C*, Brain MRI of patient 1, demonstrating lateral ventricular dilatation with a frontal dominance, suggesting a small caudate nucleus, thin corpus callosum (not shown), and delayed myelination in the white matter.

but adrenocorticotrophic hormone (ACTH) therapy was not effective. Combinations of multiple antiepileptic drugs, such as phenobarbital (PB), carbamazepine (CBZ), clobazam (CLB), valproic acid (VPA), vitamin B6, zonisamide (ZNS), and potassium bromide, failed to stop his seizures, and his development was profoundly delayed. He had not acquired eye contact or head control by the age of 18 mo. Consecutive brain magnetic resonance imaging (MRI) revealed progressive brain atrophy (fig. 1C). Patient 2 was the second child of unrelated healthy parents, and his elder sister was healthy. He was delivered spontaneously

at term, without asphyxia. After birth, he showed blinking of the left eyelid and abrupt flexion of the right arm and left leg, suggesting seizures. At the age of 1 wk, he had brief tonic seizures in clusters, with a suppression-burst pattern on EEG (fig. 2B). His physical findings were normal except for a small penis at the age of 1 mo. His routine blood tests showed normal results except for a mild elevation in liver transaminase (aspartate aminotransferase [AST] 82 IU/liter, alanine aminotransferase [ALT] 34 IU/liter). His brain MRI scan demonstrated a defect of the cavum septi pellucidi and hypoplastic corpus callosum

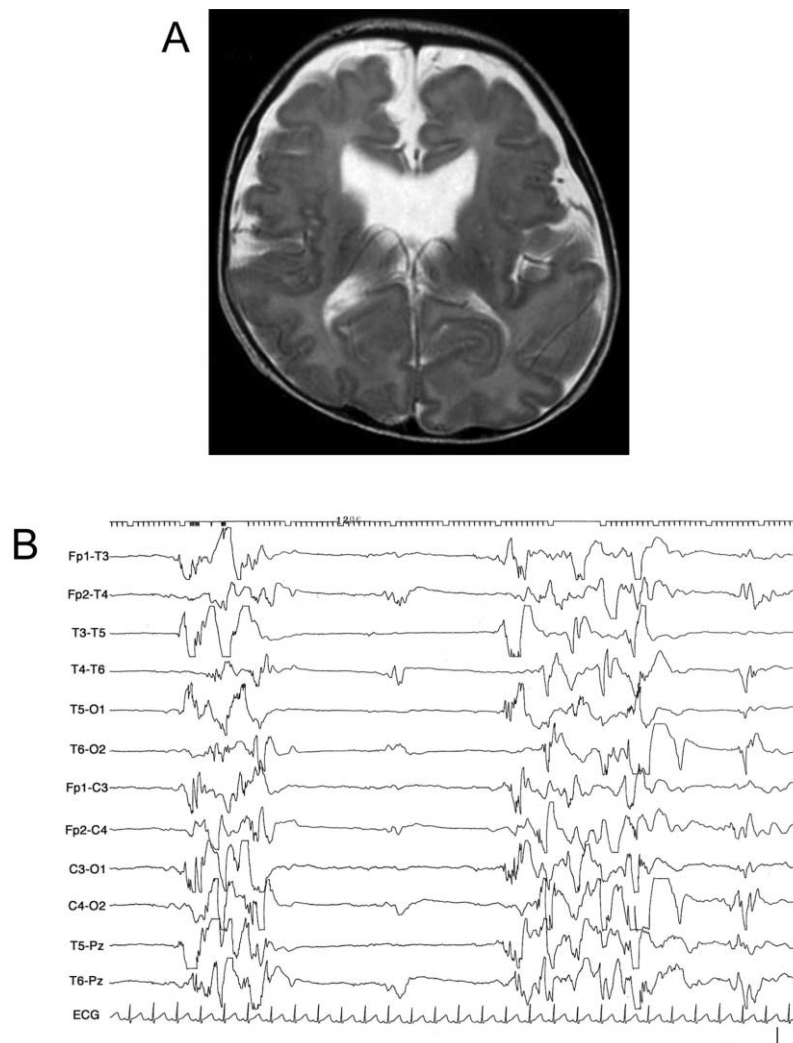


Figure 2. *A*, Brain MRI of patient 2 at the age of 8 WK shows dilatation of the anterior horns of the lateral ventricle and apparently normal gyri. *B*, Interictal EEG of patient 2 shows a marked depression of background activity, intermixed with asymmetric paroxysmal bursts.

(fig. 2A). VPA, ZNS, PB, and clonazepam were given, but his EEG at the age of 7 mo showed transformation to hypsarrhythmia. With a diagnosis of West syndrome, he received ACTH injections, with no effect. His development was severely delayed.

Blood samples were collected from three patients with sporadic EIEE and from their parents, after both informed consent from the parents and approval from the Yamagata University School of Medicine Institutional Review Board were obtained. Genomic DNA was extracted according to standard procedures and was amplified for all coding exons and flanking introns of ARX. The primers and PCR conditions have been described elsewhere.¹⁰ Direct sequence analysis of the PCR product of the first half of exon 2 was performed on an automated DNA sequencer (ABI 310 [Applied Biosystems]). Other products were analyzed by denaturing high-performance liquid chroma-

tography (DHPLC) performed using the Transgenomic WAVE system (Transgenomic).

A hemizygous 33-bp duplication in exon 2, 298_330dupGCGGCA(GCG)₉, which is thought to expand the original 16 alanine residues to 27 alanine residues (A110_A111insAAAAAAAAAAAAA) in the first polyaniline tract of the ARX protein, was found in patient 1 (fig. 3). Interestingly, the same mutation of the ARX gene was also found in patient 2. No mutation was found in their parents, suggesting that these mutations are in fact *de novo*. The mothers, however, can still be gonadal mosaics, because gonadal or somatic mosaicism was documented for the ARX gene.¹² Mutation screening of ARX showed normal results in another patient.

ARX comprises five exons encoding a protein of 562 aa, including a paired-class homeodomain and four repeats of 7–16 alanine residues called “polyalanine tracts.”^{5,9} In

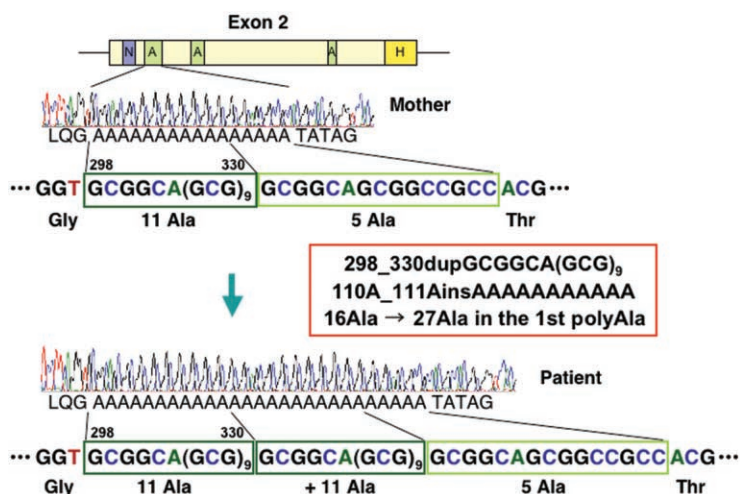


Figure 3. The structure of exon 2 of the *ARX* gene and representative electropherograms in patient 1 and his parents. Both parents have the wild-type sequence encoding 16 alanine residues, whereas the patient has a hemizygous mutant sequence with a duplication of 33 bp, which encodes 11 alanine residues and results in an expanded repeat of 27 alanine residues. Polyalanine tract (A), nuclear localization sequence (N), and homeobox (H) are illustrated in the structure of exon 2.⁹

nature, the maximum length of alanine repeats is 20.¹³ Mutation of the second polyalanine tract with eight expanded alanine residues, which is the most common mutation in *ARX*, results in nonsyndromic mental retardation; Partington syndrome (PRTS [MIM 309510]), which is characterized by mental retardation and dystonic movements of the hands; and West syndrome or infantile spasms in males.^{6,14} An expansion of 1–3 alanine residues in the first polyalanine tract containing the original 16 alanine residues has been reported in patients with mental retardation, though the expansion of 1 alanine residue has been found in 1 of 188 control males.^{15,16} An expansion of seven alanine residues in the first polyalanine tract causes West syndrome more severe than that caused by the second polyalanine tract expansion mutation of the *ARX* gene.^{14,17} So far, nine genes with expansion of the polyalanine tract have been found to be responsible for human diseases, and clinical observations demonstrate a correlation between the length of the repeat and the severity of the clinical phenotype and penetrance in *HOXD13* for synpolydactyly (SPD1 [MIM 186000]),¹⁸ *PABPN1* for oculopharyngeal muscular dystrophy (OPMD [MIM 164300]),¹⁹ and *PHOX2B* for congenital central hypoventilation syndrome (CCHS [MIM 209880]).²⁰ Longer expansion of the polyalanine tract causes a more severe phenotype, as well as other diseases caused by triplet repeat expansion, such as fragile X syndrome (MIM 300624) by *FMR1* mutation, Huntington disease (HD [MIM 143100]) by *IT15* mutation, and so on.²¹ As seen in both of our patients, the onset of epileptic seizures in EIEE is usually within the 1st mo after birth, which is distinctly earlier than 3–9 mo of age as seen in West syndrome, whereas the development outcome and life expectancy in EIEE are worse than those in West syndrome.⁴ Our obser-

vation that EIEE had longer expansion of the polyalanine tract than West syndrome may explain why EIEE starts earlier and is more severe than West syndrome.

Although the function of polyalanine tracts remains unknown, repeats of several alanine residues are frequent in transcription factors and other proteins with nuclear localization.¹³ Extended alanine repeats of *Hoxd13*, *Hoxa13*, *Runx2*, and *Sox3* spontaneously form aggregates in the cytoplasm.²² The smallest alanine-repeat expansion (+7) of *Hoxd13* was predominantly located in the nucleus and was well tolerated by the cells, whereas long expansions ($\geq +10$ alanine residues) were cytotoxic and mainly located in the cytoplasm.²³ Therefore, the size of the alanine expansions correlates with the severity of the cytoplasmic aggregation. In vitro transfection experiments of the polyalanine-expanded constructs of *Arx* cause nuclear aggregation, filamentous nuclear inclusions, and an increase in cell death, whereas cortical neurons transfected with expanded polyalanine using whole-brain electroporation form neuronal nuclear inclusions in vivo.²⁴ Recently, the human *ARX* protein was shown to be a potent transcriptional repressor.²⁵ Although the domain encompassing polyalanine tracts 1 and 2 of *ARX* seems not to be significantly contributing to the repression activity, the expansions of either polyalanine tract 1 or 2 enhance transcriptional repression activity in a manner dependent on the length of the alanine expansion.²⁵ These experimental data suggest that the expansion of *ARX* polyalanine tracts is harmful to neurons in a size-dependent manner.

Phenotypes caused by the *ARX* mutation are largely classified into two groups, a malformation group and a non-malformation group.¹⁰ Premature termination mutations and missense mutations within the homeobox of the *ARX* gene mainly cause malformation of the brain and external

genitalia, such as X-linked lissencephaly with abnormal genitalia or hydranencephaly with abnormal genitalia (XLAG [MIM 300215]) and Proud syndrome (MIM 300004), which consists of X-linked mental retardation, agenesis of the corpus callosum, and abnormal genitalia.¹⁰ Missense mutations outside the homeobox and expansions of polyalanine tracts lead to the nonmalformation group, as mentioned above.^{5,15} No apparent brain or genital malformations have been reported in patients with expansion of the polyalanine tracts, except for a boy showing transsphenoidal encephalocele and agenesis of the corpus callosum in a family affected with X-linked mental retardation²⁶; however, both our patients with an expansion of 11 alanine residues in the first polyalanine tract showed micropenis. Among patients with XLAG, missense mutations within the homeobox of *ARX* lead to less-severe genital malformation than premature termination mutations, which are thought to cause loss of function.¹⁰ Another patient without the *ARX* mutation in this report did not have genital malformation. Similar genital findings between a malformation group and EIEE caused by the longest expansion of the polyalanine tract so far reported in *ARX* bridges the gap between the malformation and nonmalformation groups and suggests a common pathological mechanism between them.

Suppression-burst pattern has also been reported in a patient with XLAG caused by a premature termination mutation in *ARX*.²⁷ The neuropathology of XLAG and *Arx*-knockout mice indicates a defect in GABAergic interneurons.^{9,11} Interestingly, an absence of GABA in the cerebrospinal fluid has been reported in a girl with EIEE and migrational disorder.²⁸ We hypothesize that an error of GABAergic system is critical to the suppression-burst pattern on EEG in patients with EIEE via dysfunction of interneurons due to the *ARX* mutation. EIEE could be designated "interneuronopathy" as well as West syndrome.⁷

The sequence of polyalanine expansions we found is an imperfect repeat of GCG including a triplet of GCA, 298_330dupGCGGCA(GCG)_n. The first 11 bp (GCGGCA-GCGGC) recurrently appear just after the sequence of duplication from nucleotides 331 to 341. This suggests a predisposition to recombination between the repeats and that unequal crossing-over could be a mutational mechanism of the expansion, as suggested for the second polyalanine expansion of *ARX* and other polyalanine expansions.^{6,29}

Knowledge of the genetic background of epilepsy is expanding, and, to our knowledge, this is the first report to reveal the genetic cause of EIEE. Mutation screening of *ARX* should be considered in patients with EIEE and in other patients showing suppression burst on EEG or infantile seizures with developmental delay.

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Web Resources

Accession numbers and URLs for data presented herein are as follows:

GenBank, <http://www.ncbi.nlm.nih.gov/Genbank/> (for *ARX* [accession number NM_139058])
 Online Mendelian Inheritance in Man (OMIM), <http://www.ncbi.nlm.nih.gov/Omim/> (for ISSX, PRTS, SPD1, OPMD, CCHS, fragile X syndrome, HD, XLAG, Proud syndrome)

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