Atypical case of Aicardi-Goutières syndrome with late-onset myoclonic status

Andrea Berger1,2, Christiane Schroeter1, Adelheid Wiemer-Kruel1, Karl Strobl1, Georg F. Hoffmann4, Dietz Rating2, Pierre Lebon3, Jan-Peter Ernst1, Nicole I. Wolf2

1 Centre of Epileptology, Paediatric Neurology, Kehl-Kork, Germany 
2 University Children’s Hospital, Paediatric Neurology, INF 155, D-69120, Heidelberg, Germany 
3 Université René Descartes, Hôpital Cochin-St. Vincent de Paul, Service de Virologie, Paris, France 
4 University Children’s Hospital, General Paediatrics, Heidelberg, Germany 

ABSTRACT – Aicardi-Goutières syndrome (AGS) is a rare, progressive, autosomal recessive encephalopathy characterised by basal ganglia calcifications, chronic CSF lymphocytosis, and negative serological investigations for the common prenatal infections. The clinical profile is characterised by acquired microcephaly, mild to severe cognitive delay and dystonia. Epilepsy is usually not prominent. We report on a 19-year-old patient with an atypical clinical course, characterized by a relatively benign presentation at onset. Epilepsy with complex-focal seizures, possibly with a visual aura and sometimes with secondary generalization, started at the age of nine years. Clinical deterioration occurred later, and at the age of 17 years he experienced severe, generalized, myoclonic attacks lasting hours, which were partly controlled by the administration of piracetam.

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showed slightly enlarged ventricles, abnormal white matter and basal ganglia. MRI at age nine years showed multiple calcifications in the subcor-tical white matter and basal ganglia. The microangiopathy is also likely caused by another characteristic feature of AGS, acrodermatitis presenting as "chilblain-like puffy swelling" (Tolmie et al. 1995, Stephenson 2002), which affects fingers, toes, ears and/or elbows. The pathogenesis of AGS is still incompletely understood. Transgenic mice with chronic overproduction of α-IFN show neuropathological abnormalities mimicking those found in AGS (Campbell et al. 1999). Skin lesions resembling AGS acrodermatitis have been reported during treatment with α-IFN (Bachmeyer et al. 1996, Campo-Vogeli et al. 1998). Chronic overproduction of IFN-alpha in AGS might thus be causally related to the microangiopathy found in brain and in skin (Barth et al. 1999).

We present the atypical clinical course of a 19 year old patient fulfilling the criteria of AGS. Diagnosis was only made at age 17 years. In addition to the well-known features of AGS, the patient developed long-lasting myoclonic attacks which were difficult to treat, but responded to piracetam.

**Case report**

This 19-year-old patient was referred to our epilepsy centre for refractory myoclonic attacks that first manifested at the age of 17 years. He was the second son of non-consanguineous German parents. An older brother aged 22 was in good health.

Pregnancy and delivery were uneventful. During the first days of life, he showed cyanosis of the lips and generalized hypotonia. Bacteriological and viral (TORCH) tests of blood and cerebrospinal fluid (CSF) were negative. CSF contained 344 cells/mm³, an elevated protein level (166 mg/dL), and normal glucose. After seven days, the CSF cell count normalized but protein remained elevated at 138 mg/dL. Cerebral ultrasound, head circumference and EEG were normal at that time. Developmental milestones were mildly delayed: he sat at nine months, walked at 18 months and spoke his first words at two years of age.

At age six he was able to ride a bike but spoke only four to five word sentences. His IQ was 50. Recurrent infections (parotitis, panaurititis, lymphadenitis) occurred from age one to school age. Physical examination at the age of seven years showed multiple enlarged cervical lymph nodes and neurodermitis. Tendon reflexes were brisk with negative plantar responses. Brain CT scan at the age of seven years showed multiple calcifications in the subcor-tical white matter and basal ganglia. MRI at age nine years showed slightly enlarged ventricles, abnormal white matter signal on T2 and FLAIR sequences, and contrast enhancement in the white matter of the medial temporal region and hippocampus. The white matter changes were interpreted as microinfarctions due to a heterozygous Factor V Leiden mutation, and antithrombolytic therapy was initiated with fraxiparin.

When he was nine years old, the first seizure-suspicious episode occurred. The boy suddenly complained of headache followed by a visual hallucination of sunlight and then went to sleep. After a few hours he awoke disoriented. He clung to his father, his body stiffened and left-beating nystagmus was observed. Further episodes recurred, four and later six times a year: The boy said “Mom, I am going to have a seizure”, then, his body stiffened and head and eyes deviated to the left with clonus of the left limbs, sometimes propagating to the right. These events lasted up to 30 minutes and were often followed by deep somnolence. Interictal EEGs showed diffuse slowing without focal or epileptic changes. At age 12, the boy became increasingly disoriented. He had abrupt mood changes with depressive and agitated outbursts. Over the course of one year, he had three episodes of tonic and myoclonic spasms of one or both legs. From age 12 to 16, the patient was treated with valproic acid, carbamaze-pine, clobazam, oxcarbazepine, lamotrigine, phenytoin, phenobarbital, and sulthaine without any effect. The frequency of seizures increased to up to four or five a day associated with retrobulbar pain, shouting and left-sided or generalised myoclonus. At age 17 years, the patient was admitted to our centre for recurrent episodes of generalized myoclonus lasting hours, accompanied by sweating and tachycardia, often without alteration of consciousness. During these episodes, he was not able to eat or drink. He had lost 4 kg over the preceding months and required a gastrostomy for feeding. General medical examination showed swollen neck and axillary lymph nodes. There were no signs of acrodermatitis. Head circumference was 54 cm (25th percentile). Neurological examination showed marked spastic/dystonic quadriaparesis. Routine EEG showed continuous generalized slowing in the theta range without epileptic activity. EEG during the myoclonic episodes was contaminated by muscle artefact but did not appear epileptic. Back averaging was not done.

To improve the myoclonus, we started treatment with piracetam. The myoclonic attacks decreased significantly and the patient was able to eat and drink and sit in a chair. He drew simple pictures. Folinic acid, 4 mg/kg/day, was added to the piracetam without any noticeable clinical effect and was discontinued three months later. However, his cognitive status continued to decline. At age 19 years, while still on piracetam, he had further episodes of generalized myoclonus over a fortnight, which resolved spontaneously.
Cranial CT scan revealed extensive white matter calcification and calcifications in the basal ganglia and cortex (figure 1). MRI showed progressive supratentorial atrophy and abnormal white matter signal (figure 2). Extensive metabolic and immunological tests at different hospitals, including a muscle biopsy with assessment of respiratory chain enzymes, were unrevealing. At age 17 years, the CSF was acellular with normal protein, glucose and lactate levels. Neopterin was extremely high at 258 nM/L (normal 10-31 nM/L) and 7,8-dihydrobiopterin 26 nM/L (normal < 18 nM/L), which is compatible with an inflammatory process. 5-methyltetrahydrofolate was 2 at the lower end of the normal range (26-118 nM/L) at 8 nM/L. IFN-alpha was not detectable in the CSF, but was elevated in serum at 3 IU/mL and 2 IU/mL one year later (normal < 2 IU/mL).

Discussion

Although microcephaly and acrodermatitis were absent in our patient, the picture of progressive psychomotor deterioration, intracranial calcifications and white matter changes along with elevated serum IFN-alpha (> 2 IU/mL) and CSF lymphocytosis in the newborn period, meet the criteria of Aicardi-Goutières syndrome. At age 17, serum IFN-alpha in our patient was still elevated, although most AGS patients show normalisation of CSF IFN-alpha with age, as well as a decline in lymphocytic pleocytosis (Goutières et al. 1998). An apparent susceptibility to infection, as found in our patient’s history, has not been reported in AGS, but has been in Cree encephalitis (Black et al. 1988), which is allelic with AGS 1 (Crow et al. 2003).

The elevation of CSF neopterin is probably secondary to IFN-alpha activation and has also been described in other patients with AGS (Blau et al. 2003). Altogether, this picture is certainly compatible with AGS, but shows some atypical features – the initial presentation was relatively benign, the CSF lymphocytosis resolved quickly, and clinical deterioration occurred only late and might have been aggravated by the increasing frequency of seizures.

Possible differential diagnoses of AGS include mainly disorders with intracerebral calcifications. There was no evidence of pre- or perinatal infections including CMV, toxoplasmosis, rubella, Herpes simplex and HIV; the intracranial calcifications and the progressive clinical course have to be explained by a metabolic or neurodegenerative disorder. The “pseudo-TORCH syndrome”, also known as “microcephaly – intracranial calcification syndrome” (MICS), shares many features with AGS, but affected children have microcephaly at birth (Vivarelli et al. 2001). Fahr disease, Cockayne syndrome, mitochondrial cytopathies including Leigh’s disease, Hoyeraal-
Hreidarsson syndrome, 3-hydroxyisobutyric aciduria, dihydropyridine reductase deficiency and cases of neonatal and familial systemic lupus erythematosus are known to be accompanied by intracerebral calcifications and microcephaly. All these disorders could be excluded either by appropriate testing or because of an incompatible clinical picture.

Our patient developed, at age 9, complex-focal seizures, possibly with a visual aura and sometimes with secondary generalization, which were resistant to treatment. Late in the course of his disease he also developed myoclonic status, an unusual feature in AGS. Although patients with AGS suffering from myoclonic jerks have been reported, so far no patient with myoclonic status has been described. According to the literature, epilepsy complicates the disease in about a third of the patients, but precise descriptions of epileptic syndrome or seizure semiology are missing (Aicardi and Goutières 1984, Lanzi et al. 2005, Abdel-Salam et al. 2004). In our case, the contamination of the ictal EEG by muscle artefacts made it impossible to determine whether the myoclonus originated from cortical or from subcortical structures. Although interictal EEG showed only diffuse slowing, massive brain atrophy, enlargement of the skull bones and widened CSF spaces could have prevented reliable detection of epileptic potentials by surface EEG. The beneficial effect of piracetam does suggest a cortical origin, since piracetam is known to be effective for cortical myoclonus, but is only poorly effective for myoclonus of thalamocortical and subcortical origin (Dulac et al. 1998, Genton et al. 1999, Genton and Van Vleymen 2000). Cortical calcifications, as seen in the CT scan of our patient, also argue for altered cortical excitability and cortical myoclonus.

The genetic basis of AGS has now been elucidated. Mutations in four genes have been found in patients affected with AGS: the AGS 1 gene located on chromosome 3p21 (Crow et al. 2000) is TREX1, a DNA exonuclease (Crow et al. 2006a). The AGS2 gene (locus 13q14-q21) (Ali et al. 2006), and two, further, recently identified genes (locus 11q13.2 and 19p13.3) are RNASEH2B, RNASEH2C and RNASEH2A forming the ribonuclease H2 enzymatic complex (Crow et al. 2006b). The precise disease mechanism is still not understood. Impaired function of one (or more) of the four nucleases could lead to elevated levels of endogenous DNA-RNA hybrids, possibly causing an inflammatory response with increased IFN-alpha production (Crow et al. 2006a,b). Results of mutation analysis in our patient are still pending. In spite of these recent advances, the diagnosis of the syndrome still remains a clinical one. At present, there is no effective causal therapy for AGS. It is expected that the identification of the genetic basis of AGS will clarify the relationship between the genetic and immunological abnormalities in this syndrome and provide helpful clues for future therapy.

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