Macrocephaly, epilepsy, autism, dysmorphic features, and mental retardation in two sisters: a new autosomal recessive syndrome

Karen Helene Ørstavik, Petter Strømme, Johan Ek, Ansgar Torvik, Ola H Skjældal

Abstract
We report two sisters with macrocephaly, epilepsy, and severe mental retardation. The first child was a 14 year old girl born at term after a normal pregnancy, with birth weight 3600 g and occipitofrontal circumference (OFC) 36 cm (75th centile). Her head size increased markedly during the first six months of life, and was later stable at 2-3 cm above the 97.5th centile. Her development was characterised by psychomotor delay, epilepsy, and autistic features. Her face appeared mildly dysmorphic with a large forehead, short philtrum, and bushy eyebrows. Her younger sister was also born at term with birth weight 2600 g and OFC 34 cm (25th centile). She also developed postnatal macrocephaly with OFC 2 cm above the 97.5th centile and the same mild dysmorphic facial features as her sister. Her development was also characterised by psychomotor delay, autistic features, and epilepsy. In addition, she suffered from coeliac disease. She died unexpectedly at the age of 5 years, probably from an epileptic attack. Necropsy confirmed megalencephaly but no other pathological changes were found. The clinical features in these two sisters do not fit with any known syndrome and may represent a previously unrecognised autosomal recessive disorder.

Keywords: macrocephaly; autistic; mental retardation; autosomal recessive

Although macrocephaly is more commonly associated with mental retardation than macrocephaly, the London Dysmorphology Database has included 105 mental retardation-macrocephaly syndromes. We report here two mentally retarded sisters with macrocephaly, epilepsy, autistic features, and mild craniofacial dysmorphism. The observations do not fit with any previously reported syndrome.

Case reports
PATIENT 1
She was a 14 year old girl born at term after a normal pregnancy. Birth weight was 3600 g, length 52 cm, and OFC 36 cm (75th centile). Head size increased rapidly after birth, but after 6 months the OFC remained stable 2-3 cm above the 97.5th centile.

Developmental milestones were delayed; she sat at 12 months, walked at 28 months, and did not learn to speak coherently. Since the age of 4 she has had epileptic seizures characterised by series of eye blinking. From the age of 11 she developed generalised complex seizures with nocturnal apnoea, myoclonus, gurgling sounds, and increased salivation. EEG showed slow background rhythm, epileptogenic spike-wave activity in both temporal lobes, and outbursts of rhythmic 1-2/second delta wave activity in the right temporoparietal region accompanied by staring spells.

When last examined at the age of 14 her OFC was 60 cm (3 cm >97.5th centile), while height and weight were at the 50th and 97.5th centiles, respectively. She had mild dysmorphic features with a high and broad forehead, deep set eyes, short philtrum, thick hair, bushy eyebrows, and a hairy upper lip (fig 1). Her mood was flat and monotypic and her behaviour was bizarre and autistic-like with lack of contact and immediate and postponed echolalia. She would not initiate complicated movements like getting dressed. Neurological examination, including fundoscopy, was normal. Her IQ was estimated as 20-35.

PATIENT 2
She was the younger sister of patient 1 who was born after an uncomplicated term pregnancy. Birth weight was 2660 g, length 49 cm, and OFC 34 cm (25th centile). She also developed postnatal macrocephaly with OFC 2 cm above the 97.5th centile from the age of 6 months. Her development was delayed. She walked at the age of 24 months and exhibited the same autistic behaviour as her older sister. From the age of 5 months she started to have partial epileptic seizures with episodes of apnoea followed by drowsiness. At the age of 2 years she developed generalised complex seizures with opisthotonus, teeth grinding, fothing at the mouth, eye blinking, and myoclonic movements. The seizures were predominantly nocturnal and frequently occurred at the time of awakening or falling asleep. Intercital EEG showed generalised high voltage dysrhythmia, while ictal EEG showed a generalised 6/second spike-wave activity, predominantly in the frontal regions.

From the age of 2 she developed clinical signs of malabsorption and coeliac disease was diagnosed. At the age of 4 the OFC was 56 cm (2 cm >97.5th centile) and her height was at
the 50th centile. Like her older sister she also had a prominent forehead and a short philtrum (fig 2).

She died unexpectedly at the age of 5 years, probably because of a nocturnal epileptic attack, as she had froth around her mouth when she was found dead in the morning. At necropsy all parts of the brain had normal proportions and were histologically normal. The brain weighed 1815 g. The mean brain weight at the age of 5 is 1200 g, and the neuropathological examination thus confirmed that the macrocephaly was the result of megalencephaly. The spinal cord and visceral organs were normal on gross and microscopic examination. The intestines were not examined histologically.

FAMILY HISTORY
A younger sister was healthy. The parents were healthy and unrelated and of normal intelligence. The OFC of the mother was 57.5 cm (97th centile) and of the father 61 cm (97th centile). Their head size at birth and the postnatal rate of brain growth was unknown. There was no history of mental retardation or epilepsy in the family. The father had a healthy brother of normal intelligence with OFC 59 cm (90th centile).

Supplementary investigations
Cranial x rays showed an enlarged neurocranium in all three dimensions of both patients. Otherwise the examination was unremarkable. Cerebral CT and MRI showed normal ventricular size and normal brain parenchyma, including that of the basal ganglia and cerebellum. Light and electron microscopic examination of a muscle biopsy was normal. A biopsy from the small intestine of patient 1 was also normal. Her spinal fluid had a normal protein content with normal protein electrophoresis. Urinary gas chromatography and mass spectroscopy did not show any metabolic abnormalities. Since macrocephaly and epilepsy are features of glutaric aciduria type 1, the activity of glutaryl-CoA-dehydrogenase in cultured fibroblasts from patient 1 was examined and found to be normal. Both sisters had a normal female karyotype. Screening for fragile X in 50 cells examined was negative.

Discussion
The main characteristics of these two sisters were early postnatal macrocephaly, epilepsy, peculiar craniofacial features, autistic behaviour, and mental retardation. The large heads of our patients could be part of a dominantly inherited benign macrocephaly unrelated to their mental and neurological dysfunction. However, the early development of pronounced macrocephaly with a high and broad forehead was more likely to be related to a brain disorder. They had no loss of abilities and the disorder therefore did not seem to be progressive. We found no biological marker suggestive of degenerative or metabolic disease.

Figure 1  Patient 1 aged 5 years (A) and 14 years (B). Note bushy eyebrows and blank, staring facial expression.

Figure 2  Patient 2 aged 2 years. Note large forehead.
and, except for macrocephaly, the postmortem examination in patient 2 was normal. This girl probably died from an epileptic attack (SUD). A predisposing factor for SUD in this case may have been the tendency for epileptic ictal apnoea. An association between epilepsy and coeliac disease has been established. However, since the older sister did not have coeliac disease, it is most likely that her gluten enteropathy was not related to the neurological abnormalities.

Several sibships have been reported with a mental retardation-macrocephaly syndrome where healthy family members had large heads. Fryns et al reported a family with three mentally retarded macrocephalic sisters. Their mother and paternal grandfather also had large heads but normal intelligence. The authors concluded that this was probably a dominantly inherited macrocephaly-mental retardation syndrome. However, these patients had short stature and very coarse features which were different from the dysmorphic features in our patients. Cole and Hughes reported six probands with macrocephaly and developmental delay and a previous diagnosis of Sotos syndrome. These patients were all very similar and all of them had one parent with a large head. However, they seemed less neurologically impaired than our patients.

Sotos syndrome and some other macrocephaly-mental retardation syndromes include increased height and general overgrowth in infancy and childhood. Our patients were of normal height.

Buttiens et al described a possible new autosomal recessive macrocephaly-mental retardation syndrome in two sisters and a brother. The sibs had developed postnatal macrocephaly but had additional findings, including coarse facial features and ocular abnormalities with severe myopia, not present in our patients. Fryns et al reported a sister and brother who had some facial resemblance to our two sisters. These sibs also had macrocephaly with a high and broad forehead, deep set eyes, a short philtrum, and epilepsy. However, in addition they had a progressive spastic paraparesis, and they were less retarded than our patients.

In our family both parents had large heads. It cannot be excluded that the two sisters were homozygous for a gene causing borderline macrocephaly in the parents. However, this possibility seems unlikely, since both parents were of normal intelligence and their facial features did not resemble those of their offspring. The syndrome boundaries of the reported macrocephaly-mental retardation syndromes are poorly delineated. To our knowledge the findings in these two sisters do not fit sufficiently with previously reported syndromes and may represent a new autosomal recessive mental retardation-macrocephaly syndrome.