A newborn girl with suspected Silver-Russell syndrome
We present the case of a girl with suspected Silver-Russell syndrome (SRS). SRS is a rare, clinically and genetically heterogeneous disorder involving intrauterine and postnatal growth retardation with a wide spectrum of additional dysmorphic features. Because of the clinical heterogeneity, diagnosis can be rather subjective and is not always easy to confirm.

The girl, third child of a 30-year-old mother, was born prematurely at 31 6/7 weeks of gestation by Cesarean section because of a pathologic CTG. Intrauterine growth retardation had been diagnosed at 26 6/7 weeks of gestation. At that time, amniocentesis and TORCH serologies were performed with inconspicuous results.

After delivery, the girl was cyanotic, bradycardic with no spontaneous respiration. Bag and mask ventilation was needed during the first minute of life. Apgar scores were 3, 9 and 8 at 1, 5 and 10 minutes, respectively. Birth weight was 980 g (P<3), length 32.5 cm (P<3) and head circumference 29.5 cm (P25-50). Besides a disproportionately large appearing head and a high forehead, downturned corners of the mouth (Fig. 1) and an asymmetry of the limbs (Fig. 2) were striking. Postnatal growth was poor (Fig. 3 and 4).

Because of these clinical findings the diagnosis of SRS was suspected and a molecular analysis was initiated,
but showed no abnormalities. The suspected diagnosis of SRS in our case therefore remained based solely on clinical criteria and could not be confirmed genetically.
Large appearing head, high forehead, downturn corners of the mouth.
Asymmetry of the legs.
Weight and height (blue: prenatal, red: postnatal measurements).
Head circumference (blue: prenatal, red: postnatal measurements).
Silver-Russell syndrome (SRS) was originally described by Silver and colleagues in 1953 (1) and little later by Russell in 1954 (2). They described children with characteristic facies, low birthweight, body asymmetry and growth retardation. Today, SRS is seen as a clinically and genetically heterogeneous disorder involving intrauterine and postnatal growth retardation with a wide spectrum of additional dysmorphic features (3).

Because the syndrome is not only clinically but also genetically heterogeneous (4), diagnosis is not always easy to confirm. Only about 10% of affected individuals have maternal uniparental disomy (UPD) of chromosome 7 (inherting 2 copies of maternal chromosome 7, with no paternal contribution) (5), a defect which may be documented by molecular analysis. If no UPD can be found, like in our case, the diagnosis remains based solely on clinical criteria. In 1999 Price proposed such diagnostic criteria for SRS (6):

1. Birth weight more or equal to 2 SD below the mean
2. Poor postnatal growth more or equal to 2 SD below the mean at diagnosis
3. Preservation of occipitofrontal head circumference
4. Classic facial phenotype (small triangular face)
5. Asymmetry (especially of the limbs)

Growth failure is the primary abnormality (7). Infants present with intrauterine growth retardation, feeding difficulty, failure to thrive, excessive sweating, a
tendency towards fasting hypoglycemia or postnatal growth retardation. Older children and adults do not manifest clinical features as clearly as infants or young children. Intelligence may be normal or there may be learning disabilities (8).

Although growth hormone (GH) levels usually are normal in affected individuals, children with SRS exhibit abnormalities of pulsatile growth hormone secretion and GH treatment may have a positive effect on the growth pattern (9). Long-term studies have not yet been performed on a large number of patients. The eventual outcome and possible adverse effects of GH treatment therefore have to be further investigated before GH therapy can be introduced as standard treatment in SRS.


