Case Report

Torticollis as the Main Presentation in a Child with Russell-Silver Syndrome: A Case Report

Mohsen Javadzadeh,1 Hedieh Saneifard,2 and Amir Hossein Hosseini2

1 Imam Hossein Hospital, Shahid Beheshti University of Medical Sciences, Tehran, Iran
2 Tajrish Hospital, Shahid Beheshti University of Medical Sciences, Tehran, Iran

Correspondence should be addressed to Amir Hossein Hosseini, amir1981hosseini@gmail.com

Received 16 September 2012; Accepted 3 October 2012

1. Introduction

Russell-Silver syndrome (RSS) was first described in 1953; it is a genetically heterogeneous imprinting syndrome that affects approximately 1/50000 children [1]. This syndrome is characterized by severe intrauterine growth retardation (IUGR) accompanied by postnatal growth deficiency. The birth weight of affected infants is typically two or more standard deviations (SDs) below the mean, and postnatal growth is two or more SD below the mean for length or height [1].

Affected individuals typically have proportionately short stature, with normal head circumference. Some other features which are less common are fifth-finger clinodactyly, typical facial features with triangular faces characterized by broad forehead and narrow chin, and limb-length discrepancy that may result from hemihypertrophy with diminished growth of the affected side [2].

Several helpful diagnostic scoring systems have been developed for RSS, and the more recent studies of Netchine et al. [3] and Bartholdi et al. [4] focused primarily on phenotypic findings. Nonetheless, many persons with RSS lack typical clinical features and have a more subtle presentation that is not consistent in all patients. For these atypical cases who lack the cardinal features of this syndrome and the clinical diagnosis that may not be made with certainty, genetic tests may be used to help to confirm the diagnosis in a significant percent of patients. Though the above tests help in confirming the diagnosis in doubtful cases, there are no tests to definitely rule out the diagnosis [3, 4].

We describe a 10-year-old girl with all of the main clinical features of RSS. Along with the typical clinical findings, she had torticollis, which has not been described before in this syndrome even among the minor features, let alone as the presenting complaint.

2. Case

A 10-year-old girl was admitted to our hospital with the chief complaints of progressive torticollis and limping. Parents had noticed the torticollis since a few weeks earlier, but they reported that limping had begun since 3 years earlier and had become worse throughout the years. Conception was natural and the pregnancy was uncomplicated. Patient was born on 39 weeks of gestation, by caesarian section. Fetal ultrasound was reported as normal in the first and the second trimesters. IUGR and oligohydramnios were detected only one week prior to birth. Birth weight was 2250 g (<10th weight for
gestational age percentile). Apgar score was 8 at 5 and 9 at 10
minutes. There were no respiratory, neurologic, metabolic,
or infectious complications at neonatal and infantile period.
The patient and her parents denied the history of any drug
consumption or abuse. She suffered from failure to thrive
with sparing of her head circumference. Her neurological
development had been within normal limits initially but
the parents reported that their daughter had become less
and less communicative gradually over the years. She had
developed a slurred speech and recently had much difficulty
in performing the school tasks. She also suffered from
limping which the parents thought to be because of right
lower limb weakness. Due to poor control over right hand,
especially during writing, she had become an obligatorily left
handed. Two weeks before her admission in our hospital,
she developed an acute-onset, nontraumatic left-sided neck
development. On admission, physical examination revealed a
girl with a weight of 19 kg (standard deviation score (SDS) =
−3.7), a height of 125 cm (SDS = −2.6), head circumference
of 57 cm (within normal range for age and gender). No
abnormalities in vital signs were detected. She was conscious
and generally cooperative during the examination. Other
remarkable findings were triangular face, torticollis and
facial asymmetry (Figure 1), leg length discrepancy, and right
hemihypertrophy. Chewing was disturbed due to right facial
hemi hypertrophy. Kyphoscoliosis and asymmetrical lower
extremities were also observed (Figures 2, 3, and 4). Parents’
examination about syndromic face or posture and torticollis
was not significant. There was a normal range of movement
in all the extremities, but mild hypotonia, especially of the
legs, was observed. In accordance to Tanner staging sexual
maturity score, she was considered B3P2.

Neurologic examinations, including cerebellar tests, were
unremarkable. Psychiatric tests were also not significant
except that she seemed shy with a low self-confidence.
Laboratory examination revealed a normal complete
blood cell count, biochemistry, liver and thyroid function
tests, and venous blood gas analysis. Rheumatologic and
metabolic investigation of serum, cerebrospinal fluid, and
urine samples showed no remarkable abnormality.

Neuroimaging and neuromuscular studies were performed;
brain MRI, MRV, MRA, and cervical MRI, in addition to electromyography and nerve conduction velocity,
produced no positive yields. Ophthalmologic exam by
pediatric ophthalmologist showed no retinal changes in favor
of hereditary neurometabolic or neurodegenerative diseases.
Also no refractory error was found and a compensatory
torticollis of ocular origin was ruled out. Sonographic eval-
uation of neck structures was performed, and a remarkable
size difference of sternocleidomastoid (SCM) muscles was
revealed (right SCM = 11 mm, left SCM = 6 mm).

Considering the positive history of IUGR and postnatal
growth retardation and fulfilling the phenotypically features
of triangular face, short stature, normal head growth,
hemi hypertrophy, gradual regression in speech, and after
performing a thorough exclusive investigation, she was
diagnosed as Russell-Silver syndrome (RSS).
Table 1: Clinical criterion for Russell-Silver Syndrome.

| Major criteria                                                                 | Minor criteria                                                                 | Supportive criteria                      |
|--------------------------------------------------------------------------------|--------------------------------------------------------------------------------|------------------------------------------|
| (i) Intrauterine growth retardation/small for gestational age (<10th percentile) | (i) Short (arm) span with normal upper-to-lower segment ratio                   | (i) Café au lait spots or skin pigmenitary changes                                      |
| (ii) Postnatal growth with height/length <3rd percentile                      | (ii) Fifth-finger clinodactyly                                                  | (ii) Genitourinary anomalies (cryptorchidism, hypospadias)                              |
| (iii) Normal head circumference (3rd–97th percentile)                         | (iii) Triangular faces                                                          | (iii) Motor, speech, and/or cognitive delays                                           |
| (iv) Limb, body, and/or facial asymmetry                                       | (iv) Frontal bossing/prominent forehead                                         | (iv) Feeding disorder                                                           |
|                                                                                  |                                                                                  | (v) Hypoglycemia                                                                     |

3. Discussion

RSS is a clinically and genetically heterogeneous disorder. It was independently described by Silver et al. [5] who emphasized the short stature and “congenital hemi hypertrophy” of these children and by Russell [6] who focused his report on the “intrauterine dwarfism” and the “craniofacial dysostosis” associated with this syndrome. RSS is considered as an imprinting syndrome. Intrauterine and postnatal growth retardation plus variable additional features including fifth-finger clinodactyly, limb hemi-hyperplasia, a typical facial phenotype, and some learning disabilities characterize it. Despite the recent advances in the genetic characterization, the diagnosis of this syndrome still relies mainly on the clinical phenotype. As compiled from information included in the studies of Netchine et al. [3], Eggermann et al. [7], Bartholdi et al. [8], and Wakeling et al. [9], the diagnosis of RSS and supportive laboratory testing should be considered in individuals who have three major criteria or two major and two minor criteria (Table 1). Nevertheless, when doubtful about the diagnosis in a specific case based on clinical findings, genetic tests may be used to confirm the diagnosis.

The most common genetic mechanism underlying RSS, responsible for approximately half of all cases, is hypomethylation on the paternal allele of H19/IGF2, located at 11p15 [10].

Our patient fulfilled all of the four major criteria, plus one minor criterion (triangular face) and two of the supportive criteria (slurred speech and learning disabilities); so the diagnosis of RSS was clinically certain. But, torticollis, as one of our patient’s chief complaints, has not been considered as a classic or even supportive criteria in RSS [7–9].

Nontraumatic, acquired torticollis has been reported in rare syndromes. Grisel’s syndrome is a non-traumatic, fixed rotary subluxation of C1 on C2 (atlantoaxial). This syndrome usually is a postoperative complication and can result in acquired torticollis, as reported by Battiaata and Pazos [11].

During the course of systemic vasculitis different types of eye involvement and subsequent compensatory torticollis may develop. Churg-Strauss syndrome (allergic granulomatous angiitis) is one of these vasculitides [12]; but in thorough investigation no finding was detected in favor of vasculitis syndromes or eye disease in our patient.

Severe craniofacial anomalies in RSS are generally uncommon. Some individuals have Pierre Robin sequence and cleft palate. Wakeling et al. [9] found cleft palate or bifid uvula in 7% of their patients with 11p15.5 methylation defects and in no patients with maternal UPD7. Dental and oral abnormalities are rare. Microdontia, high-arched palate, and dental crowding secondary to the relative micrognathia and small mouth have been reported [9, 13–15]. RSS is a constellation of variable and sometimes new manifestations. As said earlier, ultrasonography showed that in our patient asymmetry of SCM muscles resulted in torticollis. As said in case presentation, our patient had a history of IUGR. There exist a few genetic conditions in which IUGR can also be a significant feature including Three M syndrome (3-M), Dubowitz syndrome [16] but in according to major features of these topics, our patient does not fulfill their criteria.

In conclusion, we are going to emphasize that asymmetry of the limbs is a well-known phenomenon in RSS but asymmetry of SCMs is a remarkable and novel finding.
References

[1] S. Abu-Amero, D. Monk, J. Frost, M. Preece, P. Stanier, and G. E. Moore, “The genetic aetiology of Silver-Russell syndrome,” *Journal of Medical Genetics*, vol. 45, no. 4, pp. 193–199, 2008.

[2] G. Binder, M. Begemann, T. Eggermann, and K. Kannenberg, “Silver-Russell syndrome,” *Best Practice and Research: Clinical Endocrinology and Metabolism*, vol. 25, no. 1, pp. 153–160, 2011.

[3] J. Netchine, S. Rossignol, M. N. Dufourg et al., “11p15 ICR1 imprinting center region 1 loss of methylation is a common and specific cause of typical Russell-Silver syndrome: clinical scoring system and epigenetic-phenotypic correlations,” *Journal of Clinical Endocrinology and Metabolism*, vol. 92, pp. 3148–3154, 2007.

[4] D. Bartholdi, M. Krajewska-Walasek, K. Ōunap et al., “Epigenetic mutations of the imprinted IGF2-H19 domain in Silver-Russell syndrome (SRS): results from a large cohort of patients with SRS and SRS-like phenotypes,” *Journal of Medical Genetics*, vol. 46, no. 3, pp. 192–197, 2009.

[5] H. K. Silver, W. Kiyasu, J. George et al., “Syndrome of congenital hemihypertrophy, shortness of stature, and elevated urinary gonadotropins,” *Pediatrics*, vol. 12, no. 4, pp. 368–376, 1953.

[6] A. Russell, “A syndrome of intra-uterine dwarfism recognizable at birth with cranio-facial dysostosis, disproportionately short arms, and other anomalies (5 examples),” *Proceedings of the Royal Society of Medicine*, vol. 47, no. 12, pp. 1040–1044, 1954.

[7] T. Eggermann, D. Gonzalez, S. Spengler, M. Arslan-Kirchner, G. Binder, and N. Schönherr, “Broad clinical spectrum in Silver-Russell syndrome and consequences for genetic testing in growth retardation,” *Pediatrics*, vol. 123, no. 5, pp. e929–e931, 2009.

[8] D. Bartholdi, M. Krajewska-Walasek, K. Ōunap et al., “Epigenetic mutations of the imprinted IGF2-H19 domain in Silver-Russell syndrome (SRS): results from a large cohort of patients with SRS and SRS-like phenotypes,” *Journal of Medical Genetics*, vol. 46, no. 3, pp. 192–197, 2009.

[9] E. L. Wakeling, S. Abu Amero, M. Alders et al., “Epigenotype-phenotype correlations in Silver-Russell syndrome,” *Journal of Medical Genetics*, vol. 47, no. 11, pp. 760–768, 2010.

[10] M. Chopra, D. J. Amor, L. Sutton, E. Algar, and D. Mowat, “Russell-Silver syndrome due to paternal H19/IGF2 hypomethylation in a patient conceived using intracytoplasmic sperm injection,” *Reproductive BioMedicine Online*, vol. 20, no. 6, pp. 843–847, 2010.

[11] A. P. Battista and G. Pazos, “Grisel’s syndrome: the two-hit hypothesis—a case report and literature review,” *Ear, Nose and Throat Journal*, vol. 83, no. 8, pp. 553–555, 2004.

[12] C. Vitali, F. Genovesi-Ebert, A. Romani, G. Jeracitano, and M. Nardi, “Ophthalmological and neuro-ophthalmological involvement in Churg-Strauss syndrome: a case report,” *Graefe’s Archive for Clinical and Experimental Ophthalmology*, vol. 234, no. 6, pp. 404–408, 1996.

[13] C. L. Cullen and R. K. Wesley, “Russell-Silver syndrome: microdontia and other pertinent oral findings,” *ASDC Journal of Dentistry for Children*, vol. 54, no. 3, pp. 201–204, 1987.

[14] M. L. Kulkarni, V. Venkataramana, C. Sureshkumar, and H. M. Shabeer, “Russell-Silver syndrome: a study of 3 cases,” *Annals of Dentistry*, vol. 54, no. 1–2, pp. 56–60, 1995.

[15] Z. Orbak, R. Orbak, C. Kara, and F. Kavrut, “Differences in dental and bone maturation in regions with or without hemihypertrophy in two patients with Russell-Silver syndrome,” *Journal of Pediatric Endocrinology and Metabolism*, vol. 18, no. 7, pp. 701–710, 2005.

[16] N. A. Akawi, B. R. Ali, H. Hamamy, A. Al-Hadidy, and L. Al-Gazali, “Is autosomal recessive Silver-Russel syndrome a separate entity or is it part of the 3-M syndrome spectrum?” *American Journal of Medical Genetics A*, vol. 155, no. 6, pp. 1236–1245, 2011.