Case Report

The Treatment of Flexion Myelopathy of the Thoracic Spine in Pierre Robin Sequence: A Case Report

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Background: A few studies had reported the spinal deformity associated with neurological symptoms in PRS patients. Objective: To report the management of myelopathy from flexible thoracic kyphoscoliosis in Pierre Robin sequence (PRS) patient.

Material and Method: A 15-year-old PRS patient with thoracic kyphoscoliosis developed myelopathy symptom for 4 months. Posterior instrumentation (Cotrel-Dubousset) from T1-T9 was done to correct her deformity without decompression.

Results: Her motor power improved to grade 5/5 two months after operation. Her urological symptom returned to normal at 4-month postoperatively.

Conclusion: Myelopathy from flexible kyphotic deformity in PRS patients can be successfully treated by posterior correction and fusion without decompression.

Keywords: Pierre Robin sequence (PRS), Spinal deformity, Myelopathy

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Case Report

Pierre Robin sequence (PRS) is a syndrome that associated with micrognathia, glossoptosis, and posterior U-shaped cleft palate(1). Few spinal problems were reported in PRS patients(2-4). Two reports were written, the first one was on an 8-year-old PRS patient with occipitoatlantoaxial instability and thoracolumbar scoliosis without neurological symptom(3), while the second one was on a 2-year-old PRS girl who developed progressive congenital kyphosis at cervicothoracic level with severe spinal cord compression at C7/T1 level(4). To the authors’ knowledge, a few studies had reported the spinal deformity associated with neurological symptoms in PRS patients. The objective of the present study was to report the management of myelopathy in PRS patient who had flexible thoracic kyphoscoliosis.

A 15-year-old PRS female developed thoracic kyphotic deformity four years ago (Fig. 1). She developed progressive numbness of both lower limbs for four months. She had difficulty walking and urinary incontinence. Her thoracic spine was kyphoscoliosis. Her joint laxity test(5) was negative. Neurological examination demonstrated motor strength graded 4/5 throughout her lower extremities. Slight sensory loss was found bilaterally on the whole trunk and lower extremities. The deep tendon reflexes were all hyperactive in both lower limbs with positive Babinski sign and ankle clonus. Her neurological symptoms improved after bed rest for one week.

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Fig. 1 The appearance of micrognathia in Pierre Robin sequence.
The anteroposterior and lateral radiograph revealed a marked (76°) kyphosis of T4-T8, moderate (20°) left-sided scoliosis of T4-T9, and bone maturation Risser 4 (Fig. 2). The flexion and extension X-ray were taken to assess the flexibility of curve. Her kyphosis decreased 30° in extension X-ray (Fig. 3). Her cervical spine X-ray revealed the block vertebrae at C2-3 and C4-5, no occipitoatlantoaxial instability (Fig. 4). The 3-dimension CT scan demonstrated bony dysplasia and small pedicle diameter at the level of the deformity (Fig. 5). MRI demonstrated spinal cord tethering at the apex of kyphotic curve with intra-medullary hyperintensity signal on T2W at T5-T6 levels (Fig. 6).

Posterior approach was done from T1-T9. Kyphotic deformity was corrected using hook rod system (Cotrel-Dubousset) without decompression.

Fig. 2 The anteroposterior whole spine radiograph demonstrates kyphoscoliosis of thoracic spine and bone maturation Risser 4 (A). The anteroposterior of thoracic spine revealed 20° left-sided scoliosis of T4-T9. The lateral radiograph revealed 76° kyphosis of T4-T8 (B).

Fig. 3 Preoperative lateral radiograph was taken before and after bending with fulcrum demonstrating 30° reduction of kyphosis.

Fig. 4 The lateral functional radiograph in flexion (A), neutral (B) and extension (C) reveals the symmetric block of the second and third and no atlantoaxial instability.

Fig. 5 The 3-dimension computed tomography scan showed bony dysplasia and small pedicle diameter at the level of the deformities.

Fig. 6 MRI sagittal-view T2W (A) and STIR (B) and Axial T2W demonstrated spinal cord tethering at the apex of kyphotic curve. Intra-medullary hyperintensity signal (arrow) was located at T5-T6 levels.
Local bone graft mixed with bone substitutes were used to promote fusion. The operation lasted 180 minutes. The intraoperative blood loss was 600 ml. The patient tolerated the procedure well and experienced no complications. Kyphotic correction was 30 degrees and confirmed by intraoperative fluoroscopy, postoperative X-ray and CT scan (Fig. 7).

After surgery, the patient reported improvement in her gait and numbness. Her motor power was gradually improved to grade 5/5 at 2-month post-operation. Her urological symptom returned to normal at 4-month postoperatively. She was doing well and radiography showed solid fusion at 1-year follow-up.

**Discussion**

Few studies report spinal deformity associated with PRS. Molnar et al reported an 8-year-old PRS child with a congenital kyphoscoliosis, an atlanto-occipital subluxation, and Klippel-Feil syndrome without neurological symptoms\(^\text{3}\). Kassisi et al reported a 2-year-old PRS girl who developed severe spinal cord compression at C7-T1 from progressive congenital kyphosis at cervicothoracic level\(^\text{4}\). Our PRS patient was different from the previous two cases that she did not have congenital kyphoscoliosis.

Spinal cord compression from thoracic kyphoscoliosis deformity was reported\(^\text{5-8}\). The treatment is decompression and deformity correction. Anterior decompression through right-side thoracotomy in 17-year-old boy with type 2 congenital kyphosis was performed\(^\text{5}\). Nine of 11 congenital kyphoscoliosis patients with spontaneous neurological deterioration had anterior decompression and fusion\(^\text{5}\).

However, the posterior correction, decompression, and fusion were performed with good clinical result in 84-year-old man with dynamic myelopathy\(^\text{9}\). The authors recommended posterior fusion in dynamic myelopathy. They also mentioned that decompression is not necessary in all cases\(^\text{9}\). Yamazaki et al propose that the major factors for thoracic myelopathy are kyphosis and instability\(^\text{10}\). The association between severity of kyphotic deformity and neurological symptoms was demonstrated. A 14-year-old Scheuerman’s disease patients developed cord compression from flexible kyphotic deformity that resolved after the kyphotic deformity was corrected by plaster jacket\(^\text{9}\).

In the present patient, posterior correction and fusion was chosen because there was no spinal cord compression ventrally and the curve was flexible. Her symptom also improved after one week of resting in supine position. Hook-rod construction was selected instead of pedicular screws system due to the bony dysplasia and small pedicular diameter. Therefore, we did the posterior correction and fusion from T1-T9 using CD system without decompression. The operation successfully improved her neurological problems. In the PRS patients who develop myelopathy from kyphoscoliosis, static and dynamic X-ray, CT, and MRI are recommended to determine the treatment methods. If the curve was flexible and there was no compression from disc or vertebra ventral to spinal cord, posterior correction and fusion without decompression is a safe and effective treatment.

**What is already known on this topic?**

Pierre Robin sequence (PRS) is a syndrome that associated with micrognathia, glossoptosis, and posterior U-shaped cleft palate. Some orthopedic related problems in PRS such as polydactyly, paraxial hemimelia, toe overlying, fusion of rib, and wide thoracic spinal canal are reported. There were only two cases of PRS reported occipitoatlantoaxial instability and thoracolumbar scoliosis without neurological symptoms.

**What this study adds?**

To the authors’ knowledge, this is the first report of dynamic thoracic kyphoscoliosis with spinal cord compression in PRS. We performed deformity correction by posterior fusion with instrumentation. The successful correction of kyphoscoliosis demonstrated significant spinal cord decompression. The patient’s neurological sequelae gradually resolved to normal.
Potential conflicts of interest
None.

References