Posterior-approach single-level apical spinal osteotomy in pediatric patients for severe rigid kyphoscoliosis: long-term clinical and radiological outcomes

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OBJECTIVE Spinal osteotomy in pediatric patients is challenging due to various factors. For correction of severe rigid kyphoscoliosis in children, numerous techniques with anterior or posterior or combined approaches, as well as multilevel osteotomies, have been described. These techniques are associated with prolonged operative times and large amounts of blood loss. The purpose of this study was to evaluate the clinical and radiologically confirmed efficacy of a modification of the apical spinal osteotomy (ASO) technique—posterior-only single-level asymmetric closing osteotomy—in pediatric patients with severe rigid kyphoscoliosis.

METHODS The authors performed a retrospective study of a case series involving pediatric patients with severe spinal deformity operated on by a single surgeon at a single institution over a period of approximately 5 years. The inclusion criteria were age < 14 years, rigid thoracic/thoracolumbar/lumbar kyphosis (> 70°) with or without neurological deficit and with or without scoliosis, and a minimum of 2 years of follow-up. Patients with cervical or lumbosacral kyphoscoliosis were excluded from the study. Demographic and clinical parameters, including age, sex, etiology of kyphoscoliosis, neurological examination status (Frankel grade), and visual analog scale (VAS) and Oswestry Disability Index (ODI) scores, were noted. Operative parameters (level of osteotomy, number of levels fused, duration of surgery, blood loss, and complications) were also recorded. Radiological assessment was done for preoperative and postoperative kyphosis and scoliosis as well as the final Cobb angle. Similarly, sagittal vertical axis (SVA) correction was calculated. Fusion was assessed in all patients at the final follow-up evaluation.

RESULTS A total of 26 pediatric patients (18 male and 8 female) with a mean age of 9 years met the inclusion criteria and had data available for analysis, and all of these patients had severe scoliosis as well as kyphosis. Comparison of preoperative and postoperative values showed a significant improvement (p < 0.05) in radiological, clinical, and functional parameters (Cobb angle for scoliosis and kyphosis, SVA, VAS, and ODI). With respect to kyphosis, the mean preoperative Cobb angle was 96.54°, the mean postoperative angle was 30.77°, and the mean angle at final follow-up was 34.81° (average loss of correction of 4.23°), for a final average correction of 64.15%. With respect to scoliosis, the mean preoperative Cobb angle was 52.45°, the mean postoperative angle was 15.77°, and the mean angle at final follow-up was 19.42° (average loss of correction of 3.66°), for a final average correction of 60.95%. The preoperative SVA averaged 7.6 cm; the mean SVA improved to 3.94 cm at the end of 2 years. Bony fusion was achieved in all patients. The mean number of levels fused was 5.69. The mean operative time was 243.46 minutes, with an average intraoperative blood loss of 336.92 ml. Nonneurological complications occurred in 15.39% of patients (2 dural tears, 1 superficial infection, 1 implant failure). At the 2-year follow-up, 25 of the 26 patients had maintained or improved their neurological status. One patient developed paraplegia immediately after the operation and recovered only partially.

CONCLUSIONS Analysis of data from this series of 26 cases indicates that this posterior-approach single-level technique is effective for the correction of severe rigid kyphoscoliosis in pediatric patients, providing good clinical and radiological results in most cases.


KEYWORDS kyphosis; scoliosis; deformity correction; osteotomy; posterior approach; spine

ABBREVIATIONS AP = anteroposterior; ASO = apical spinal osteotomy; COWO = closing-opening wedge osteotomy; CWO = closing wedge osteotomy; MEP = motor evoked potential; ODI = Oswestry Disability Index; PSO = pedicle subtraction osteotomy; SPO = Smith-Peterson osteotomy; SSEP = somatosensory evoked potential; SVA = sagittal vertical axis; VAS = visual analog scale; VCR = vertebral column resection.


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The correction of sagittal-plane deformity in pediatric patients is a challenge for the spine surgeon that has gripped the curiosity of many in the recent past.2,9 Loss of normal sagittal balance produces a posture that is at a biomechanical disadvantage and may cause spinal cord compression, giving rise to neurological symptoms. The aim of surgery is to fuse the spine in a balanced position as close to a normal configuration as possible with minimal morbidity and complications. Correction of kyphoscoliosis can be achieved by a single posterior approach with the help of closing wedge osteotomies (CWOs), such as the Smith-Peterson osteotomy (SPO), pedicle subtraction osteotomy (PSO), and vertebral column resection (VCR). However, the anatomical limitations of a vertebral body restrict CWO to about 35° lordosis.5,12,18 When sagittal imbalance exceeds 25 cm, consideration must be given to performing more than one CWO to obtain adequate correction.6 Similarly, excessive shortening is dangerous, since the spinal cord may then be too long for the shortened column and become kinked and possibly damaged. Gertzbein and Harris12 recommended correction to 30°–40°.

For the correction of severe rigid deformities through a posterior approach and osteotomy at a single level, authors have recently described modifications of conventional osteotomies like apical lordosating osteotomy,7 modified VCR,29 apical segmental resection osteotomy,10 and closing-opening wedge osteotomy (CWO).8,14 These surgical procedures have been associated with long operating hours and significant blood loss.8,14,29

Spinal osteotomy in pediatric patients is challenging due to various factors, including poor bone quality, rapid progression of the deformity (during physiologic growth spurts), lack of implants, and suboptimal implant purchase. The purpose of this study was to evaluate the clinical and radiological outcomes of a modification to the technique of apical spinal osteotomy (ASO), a posterior-only single-level osteotomy in the treatment of pediatric patients with severe rigid kyphoscoliosis, and to evaluate its clinical and radiological efficacy.

Methods

This study was designed as a retrospective analysis of cases in which patients with severe spinal deformity were treated by a single surgeon (V.K.) at our institution between November 2009 and August 2014. Additional inclusion criteria were age <14 years, rigid thoracic/thoracolumbar/lumbar kyphosis (>70°) with or without neurological deficit and with or without scoliosis, and a minimum of 2 years of follow-up. Patients with cervical or lumbosacral kyphosis were excluded. Data were collected and evaluated by an independent observer. All patients underwent comprehensive clinical and radiological evaluation, and demographic, clinical, neurological examination, and radiological data were collected preoperatively and at follow-up (immediately after surgery and at 3 months, 6 months, and 1 year postoperatively, as well as at the final follow-up visit 2 years postoperatively). The patients were evaluated by the operating surgeon and a junior spine surgeon; data were collected by 2 clinical spine fellows who were not part of the operating team.

Clinical Assessment

Patient demographic data (age and sex), etiology of spinal deformity, neurological status (Frankel grade), and visual analog scale (VAS) pain scores and Oswestry Disability Index (ODI) scores were collected.

Operative Assessment

Operative parameters, including the level of osteotomy, number of levels fused, duration of surgery, intraoperative blood loss, and complications, were recorded.

Radiological Assessment

Full-length standing (with hips and knees extended) anteroposterior (AP) and lateral radiographs of the spine were available for evaluation. The level of the apical vertebra, sagittal vertical axis (SVA), and Cobb angles for kyphosis and scoliosis were documented. Fusion was assessed (using the Bridwell criteria) on follow-up radiographs by an independent observer who was not part of the surgical team. All radiological images were assessed by a clinical research fellow in our institution’s radiology department with the supervision of a consultant radiologist, neither of whom was a part of the surgical team.

Operative Technique

All patients underwent 3-column spinal osteotomy, performed at the apex of their deformity—apical spinal osteotomy (ASO)—as described below.

Patients were placed prone under general anesthesia on bolsters with padding of all bony prominences. The apex of the deformity was placed over the hinge of the operating table so that when the osteotomy was closed, the table could be adjusted from a flexed position to more neutral alignment if needed. Intraoperative multimodality neuro-monitoring (monitoring of somatosensory evoked potentials [SSEPs] and motor evoked potentials [MEPs] and electromyography) was used in all cases. A posterior midline incision was performed and strict subperiosteal dissection was carried out to avoid blood loss from the upper to the lower planned instrumented levels. Facets included in the fusion levels were excised to promote intraarticular arthrodesis. Pedicle screws were inserted using freehand technique and C-arm guidance. Titanium 3.5-, 4.5-, and 5.5-mm pediatric pedicle screws were used for this purpose. Four points of fixation were secured before vertebral resection and connected with a contoured rod on one side to provide stability.

Resection of the Vertebral Column

Resection was carried out at the apex of the deformity utilizing the lever arm to the maximum effect. Because the patients in this study all had kyphosis with scoliosis, the width of the osteotomy was broader on the convex side and posteriorly for correction of both sagittal and coronal plane deformities simultaneously. The transverse process and rib heads were removed to allow complete dissection of the lateral wall of the vertebral body (Fig. 1A). Segmen-
tal vessels in the line of dissection were ligated and cut on only one side. Nerve roots in the thoracic region were sacrificed, if necessary, although nerve roots in the lumbar region were preserved throughout the procedure. The lateral wall of the pedicle was removed piecemeal. Vertebral body resection (Fig. 1B) was performed with the help of pituitary rongeurs, a 4-mm cutting burr, and curettes initially. Dissection was carried out until the anterior wall was removed, keeping the anterior longitudinal ligament, medial wall of the pedicle, and posterior wall of the vertebral body intact. A temporary rod was fixed on the ipsilateral side, and a similar procedure was repeated on the contralateral side. At the end of this process, a bony shell remained around the dural sac circumferentially (Fig. 1C). At this stage, the posterior vertebral wall anterior to the dural sac, the medial pedicle wall, and the laminae on the dorsal aspect of the dura were removed (Fig. 1D). This step is performed when an adequate amount of the vertebral body is resected. We believe that it helps in reducing blood loss from epidural veins and protects the dural sac from injury during the resection procedure. The posterior cortex of the vertebral body was initially thinned out with a diamond burr and was then removed (pushed down) with the help of reverse angle curettes (Fig. 1E).

Deformity Correction

Deformity correction (Fig. 2) was attempted by closure of the osteotomy with sequential rod contouring. Gradual correction of the kyphosis angle was attempted at every sequence of rod bending, with extension of the operating table if needed. In correcting scoliosis, compression and shortening was done more on the convex side and was asymmetrical. Although the mean arterial pressure was adequately maintained throughout the procedure, it is particularly important at this step to maintain spinal cord perfusion. The end point of column shortening in our series was direct visualization of dural buckling, any MEP/SSEP signal change or loss, or more than 15 mm of closing of the wedge on the convex side at the posterior face of the osteotomy.

Arthrodesis

Posterior fusion was performed at all instrumented levels. The number of levels to fuse was decided intraoperatively depending upon the extent of correction achieved and the gap present anteriorly after osteotomy. If the gap was less than 10 mm and sufficient correction was achieved with bone-on-bone contact after closure of the

![FIG. 1. Intraoperative images obtained during resection of the vertebral column. A: Meticulous subperiosteal elevation off the lateral vertebral body and anterolateral dissection with protection of pleural sac. B: Decancellation of vertebral body starting from the convex side. C: Completion of decancellation and intact spinal canal giving an avascular field (roller-gauze seen passing anterior to intact bony canal enclosing the dural sac). D: Removal of posterior elements with a Kerrison rongeur and stabilization with rod on one side. E: Breaking the posterior wall of the vertebral body with reverse angled curette. Figure is available in color online only.](image1)

![FIG. 2. Intraoperative images obtained during deformity correction. Left: Before closure of osteotomy. A precontoured rod can be seen. Right: Closure of osteotomy with compression to correct deformity in both sagittal and coronal planes. Figure is available in color online only.](image2)
osteotomy, only 2 levels above and below were fused; constructs involving 3 levels above and 2 levels below or 3 levels above and below the osteotomy were also used in a few patients when needed. Considering that all patients were in the range of 6–14 years of age with a significant number of thoracic deformities, efforts were made to save as many levels as possible and thus give them a chance for better truncal height and pulmonary capacity at maturity. Anteriorly, if the residual bony defect after closing osteotomy/column shortening was less than 10 mm, autogenous cancellous bone graft (bone chips) was placed. If the anterior gap was more than 10 mm, a titanium mesh cage filled with bone graft was placed. The cage was precisely sized for the anterior gap and did not cause anterior column lengthening. Additional compression was applied over the cage to lock it into place and maintain optimum correction. Wound closure was performed in layers over a negative suction drain.

Postoperative Treatment
Patients were allowed to sit up in bed within 24 hours. They were allowed to mobilize out of bed on the 2nd postoperative day with brace support. A brace was recommended for a period of 6 weeks.

Statistical Analysis
Statistical analysis was done using chi-square and Student t-tests for assessment of significance between variables; p < 0.05 was considered significant. We engaged a biostatistician for assistance with statistical analysis. A normal distribution was obtained and parametric tests were used.

Results
A total of 32 consecutive pediatric patients (21 male and 11 female) with severe kyphoscoliosis met all the criteria other than the required length of follow-up. Three of the patients were lost to follow-up and 3 had less than the required duration of follow-up; therefore, these 6 patients were excluded from analysis. Data from 26 patients (18 male and 8 female) were thus available for analysis, and these patients formed the study group. Although the inclusion criteria specified “rigid thoracic/thoracolumbar/lumbar kyphosis (> 70°) … with or without scoliosis,” all of the patients who met the criteria did, in fact, have both kyphosis and scoliosis.

The mean duration of follow-up for these 26 patients was 28.31 months (SD 5.14 months, range 24–48 months). The patients’ mean age at the time of surgery was 9 years (SD 2.5 years, range 6–14 years).

The cause of spinal deformity varied (Table 1). Congenital kyphoscoliotic deformity was diagnosed in 8 patients, kyphoscoliosis due to post tubercular infection in 7, neurofibromatosis in 5, posttraumatic kyphosis in 3, idiopathic kyphoscoliosis in 2, and Scheuermann’s kyphosis in 1 patient.

Symptoms at presentation included mechanical back pain, cosmetic deformity of the back, and neurological symptoms. Neurological deficit was diagnosed preoperatively in 9 patients (Frankel grade A in 1 case, B in 3 cases, C in 5, and E in 17) (Table 2).

Clinical Outcome
At final follow-up, the Frankel grade improved from A to B in 1 case, from B to D in 2 cases, and to E in 1 case; in all 5 cases in which the preoperative Frankel grade was C, the grade improved to E. At the end of 2 years, all patients had maintained or improved their neurological examination status except one patient who developed paraplegia immediately after surgery and recovered only partially. One patient suffered transient left-sided L-4 weakness, which had improved to normal at the final follow-up. The final Frankel grades are shown in Table 2.

Significant improvement (p < 0.05) also occurred in pain (VAS) and functional (ODI) scores. The mean VAS score improved from the preoperative value of 5.85 (SD 1.64, range 2–8) to 3.15 (SD 1.29, range 2–6). The mean ODI score was 56.85 (SD 19, range 20–90) preoperatively and had decreased to 19 (SD 8.24, range 10–40) at final follow-up (Table 3).

When we divided the cases into 2 subgroups based on etiology of spinal deformity (congenital or acquired), we found that the mean preoperative ODI score was 39.2 for the congenital cases and 64.7 for the acquired cases (Table 4); these mean scores improved to 24.8 and 16.4, respectively. The mean VAS score was 4.2 in the congenital cases and 6.4 in the acquired cases preoperatively and improved to 3.3 and 3.2, respectively.

Operative Outcome
Single-level osteotomy was performed in all patients. Osteotomies were performed from levels T-5 to L-3 (Table 5) depending on the apex of kyphosis, most commonly at the T-12 and L-1 levels (in 13 of 26 patients). Osteotomy was done at T-10 or above in 6 (23.1%) of 26 patients and at T-11 or below in 20 (76.9%) of 26 patients. The mean num-

<table>
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<th>TABLE 1. Distribution of etiologies</th>
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<td>Etiology</td>
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<tr>
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</tr>
<tr>
<td>Congenital</td>
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<tr>
<td>Post-TB</td>
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<td>Neurofibromatosis</td>
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<tr>
<td>Posttraumatic</td>
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<td>Idiopathic</td>
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<td>Scheuermann’s kyphosis</td>
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<th>TABLE 2. Neurological examination findings: Frankel’s grading</th>
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<tr>
<td>Frankel Grade</td>
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</tr>
<tr>
<td>A</td>
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<tr>
<td>B</td>
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<tr>
<td>C</td>
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<tr>
<td>D</td>
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<tr>
<td>E</td>
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TB = tuberculosis.
Radiological Outcome

The mean preoperative kyphotic Cobb angle was 96.54° (SD 15.73°), with a range of 70°–130°. After surgery, curves averaged 30.77° (SD 10.17°, range 20°–50°), yielding a significant average correction of 68.57% (p < 0.05). At final follow-up, the mean kyphotic curve was 34.81° (SD 10.15°, range 20°–60°), indicating an average final correction of 64.15%. An average loss of 4.23° at final follow-up was documented (Table 7).

Similarly, the mean scoliotic Cobb angle improved significantly (p < 0.05) from a preoperative value of 52.54° (SD 18.13°, range 26°–88°) to 15.77° (SD 5.07°, range 10°–26°) postoperatively, with average correction of 68.79%. At last follow-up, the mean scoliosis angle was 19.42° (SD 5.31°, range 12°–30°), with a mean correction of 60.95%. An average loss of correction of 3.66° at final follow-up was noted (Table 7).

The preoperative SVA averaged 7.6 cm (SD 2 cm, range 3.5–11 cm) and improved significantly (p < 0.05) to 3.94 cm (SD 0.86 cm, range 2.5–6 cm) at the end of 2 years. Bony fusion was achieved in all patients. Of 26 patients, 9 (34.62%) exhibited grade 1 fusion, 15 (57.70%) exhibited grade 2 fusion, and 1 (3.85%) exhibited grade 3 fusion per Bridwell’s criteria of radiological fusion.

Complications

Complications occurred in 8 patients (30.77%) (Table 8). In 2 cases, dural tears were diagnosed intraoperatively and repaired primarily by suturing of the dura followed by watertight closure of the operative wound. One patient had an implant failure (screw pullout), which required revision surgery. One patient developed a superficial wound infection, which was managed conservatively with antibiotics and regular dressings. One patient (with congenital kyphoscoliosis, treated with ASO at the L-1 level) had a new-onset neurological deficit involving the left L-4 nerve root (quadriiceps weakness 3/5), and in 2 other patients the neurological examination findings worsened from preoperative Frankel grade C to A in the immediate postoperative period. One patient developed complete paraplegia (congenital kyphoscoliosis, ASO at the L-1 level), which had resolved only partially by the final follow-up evaluation. It was later realized that the osteotomy was incomplete and forced closure with load on the implants led to translation within the narrow canal. The changes in neuromonitoring signals were detected during manipulation and closure of the osteotomy. Upon realization of this, the correction maneuver was reversed with adequate ventilation, reduction of inhalational anesthetics, and maintenance of mean arterial pressure. In spite of these measures, the neuromonitoring signals did not return to baseline levels and amplitudes had increased only partially by the time closure was complete. The patient was also given a short course of intravenous steroid therapy, but only partial improvement was found in neurological status even at the final follow-up (Frankel grade D). The same patient suffered from screw pullout (as mentioned above) and construct failure with loss of correction, for which a revision surgery with extension of fixation and redirection of the loosened screw was performed.

Illustrative Case

This 14-year-old boy presented with a progressive back deformity and no neurological deficits. He had a positive sagittal balance with compensatory increased lumbar lordosis. Radiographs (Fig. 3) revealed block vertebrae T10–12 resulting in a rigid kyphotic deformity of 75°. An apical osteotomy was performed with correction of sagittal balance. The radiographs obtained 2 years postoperatively (Fig. 4) show fusion and maintained correction and sagittal balance without any implant-related complications.
Discussion

The present study analyzed the clinical and radiological efficacy of apical spinal osteotomy (ASO) performed using a posterior-only single-level technique for severe rigid kyphoscoliosis in pediatric patients. The uniqueness of the study is the rarity of the study group with diverse etiologies and severity of deformity in the pediatric population. These patients were treated with a single-level osteotomy at the apex of the curve, hence the term "apical spinal osteotomy." Rigid kyphoscoliosis is quite common in young children.2,10,14–16,23,25,29 The “pitched-forward” posture causes back muscles to strain in an effort (successful or unsuccessful) to reduce the tilt, resulting in additional energy expenditure that leads to reduced functional capacity as well as a poorer quality of life due to mechanical and neurological symptoms.3 The aim of surgery is to fuse the spine in a balanced position that is as close to a normal configuration as possible.

Anterior and Posterior Versus Posterior-Only Approach

A combined anterior release and fusion followed by posterior spinal fusion and instrumentation has often been performed for large and stiff curves.31 Both open and endoscopic approaches have shown greater negative impact on pulmonary function than the posterior-only approach.17,19 Luhmann et al.22 concluded that patients treated with pedicle screw–only instrumentation presented results similar to those for patients with combined treatment (60.7% vs 58.5%), without negative effects on pulmonary function from anterior release. With the advent of modern instrumentation and pedicle screw systems, a single posterior approach is preferable to reduce morbidity to the patients. All patients in the present series were operated on with a single posterior-only approach.

Column-Shortening Procedures

In highly rigid severe spinal deformities, conventional correction methods, such as posterior-only procedures or anterior release and posterior instrumentation, are usually unsatisfactory and a more aggressive approach is necessary.26 Closing wedge osteotomies, such as SPO, PSO, and VCR, can provide acceptable clinical and radiological results for patients with sagittal imbalance.4 Suk et al. reported on a posterior-only approach with a VCR for fixed lumbar spinal deformities26 as well as for severe rigid scoliosis.25,27 These authors reported excellent surgical correction with minimal long-term complications. However, anatomical limitations of vertebral bodies restrict CWO to about 35° lordosis.5,12,18 When sagittal imbalance exceeds 25 cm, consideration should be given to performing more than just one CWO to obtain adequate correction,9 increasing the risks and complications of the surgery. In our technique, correction of severe rigid deformities was achieved with the help of single-level apical osteotomy for all patients. The end point of column shortening in our series was direct visualization of dural buckling, any MEP/SSEP signal change or loss, or more than 15 mm of closing of the wedge on the convex side at the posterior face of the osteotomy. Also, if the residual bony defect after osteotomy closing/column shortening was less than 10 mm, autogenous cancellous bone chips were placed. If the anterior gap was more than 10 mm, a titanium mesh cage filled with bone graft was placed. The benefits of performing an apical osteotomy are efficient correction at the very site of rigid deformity, the potential to correct deformity in both sagittal and coronal planes, and direct bone-to-bone contact instead of spanning reconstruction, giving better fusion rates and fewer implant-related complications.

Etiology and Patient Age

Chen et al.,10 Shimode et al.,23 Bakaloudis et al.,1 and Wang et al.28 studied rigid kyphoscoliosis of varied etiologies and have documented the efficacy of thoracic PSO, VCR, and apical resection procedures with final corrections in the range of 54%–69%. The mean age of the patients in these studies was in the range of 12–34 years. Similarly, Lenke et al.21 reported the results of VCR in 35 pediatric patients with a mean follow-up of 2 years. The mean age of the patients in their series was 11 years (range 2–18 years). The patients in that case series underwent either single-level (n = 20) or multilevel (n = 15) osteotomies, and the authors documented overall correction rates of 51%–60%, with an average blood loss of 691 ml. In our series, the patients’ mean age was lower (9 years, range 6–14 years), all of the patients underwent single-level ASO for spinal deformities of varied etiologies as documented above, and correction rates of 64% and 60.9% were achieved for kyphotic and scoliotic curves, respectively. The mean blood loss in our series was 337 ml.

Blood Loss

Treatment of severe kyphoscoliosis with osteotomies is associated with significant blood loss. Various studies1,8,14,23,29 on the treatment of rigid kyphoscoliosis with apical lordosating osteotomy, SWO, multilevel modified VCR, and COWO have documented blood loss in the range from 717 to 3340 ml. In their 2013 report on a multicenter trial, Lenke et al.28 emphasized the magnitude of blood

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**TABLE 7. Radiological outcomes**

<table>
<thead>
<tr>
<th>Timing &amp; Measure</th>
<th>Kyphosis</th>
<th>Scoliosis</th>
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<tbody>
<tr>
<td>Preop (°)</td>
<td>96.54</td>
<td>52.54</td>
</tr>
<tr>
<td>Postop (°)</td>
<td>30.77</td>
<td>15.77</td>
</tr>
<tr>
<td>Final follow-up (°)</td>
<td>34.81</td>
<td>19.42</td>
</tr>
<tr>
<td>Loss of correction (°)</td>
<td>4.23</td>
<td>3.66</td>
</tr>
<tr>
<td>Final correction (%)</td>
<td>64.15</td>
<td>60.95</td>
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**TABLE 8. Complications**

<table>
<thead>
<tr>
<th>Complication</th>
<th>No. of Patients (%)</th>
<th>Final Outcome</th>
</tr>
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<tbody>
<tr>
<td>Dural tear</td>
<td>2 (7.69%)</td>
<td>Healed</td>
</tr>
<tr>
<td>Implant failure</td>
<td>1 (3.85%)</td>
<td>Revision surgery</td>
</tr>
<tr>
<td>Superficial wound infection</td>
<td>1 (3.85%)</td>
<td>Healed</td>
</tr>
<tr>
<td>Neurological deficit</td>
<td>4 (15.38%)</td>
<td>Recovered in 3 cases, persistent in 1 case</td>
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loss possible in VCR. They reported an average blood loss of 1610 ml. Similarly, Sponseller et al., 24 in their study of children with neuromuscular scoliosis treated with VCR, reported a mean blood loss of 1785 ml. Our mean blood loss (337 ml, range 200–500 ml) was much less than in the previously described series. We believe that in our technique the preservation of the dural sac within the bony shell until adequate resection of the vertebral column is performed significantly prevents blood loss from the bleeding epidural veins. Also, in patients < 14 years old the tissue planes are well delineated with minimal degeneration.

Correction Magnitude

In correction of high-magnitude rigid curves, similar correction rates have been reported for both conventional (SPO, PSO, and VCR) and newer unconventional osteotomies (apical resection osteotomy, SWO, and COWO). However, conventional osteotomies may require correction at multiple levels, which increases the morbidity of surgery. Bakaloudis et al.1 reported an overall correction of 65% with PSO in pediatric patients. Chen et al.10 documented correction of 69.87% with apical resection osteotomy. In our series, the mean kyphotic curve was 96.5° preoperatively and 35° at final follow-up, indicating an average final correction of 64.15%, which remained essentially consistent at the end of 2 years (the change of 4.2° between postoperative and final measurements was not clinically significant). Our results were similar to those of many studies mentioned in the literature,1,7,10,23,28 thus proving efficacy of the technique.

Patient Satisfaction Rates

PSO and VCR have provided good results not only in radiological terms but also in clinical terms for patients with fixed sagittal imbalance caused by multiple etiologies, but only a few studies have reported functional outcomes. After performing PSO in 27 patients with fixed sagittal imbalance, Bridwell et al.6 noted that the mean ODI score improved from 51.21 preoperatively to 35.75 at the last follow-up evaluation, and 92.3% of the patients were found to be satisfied with the treatment according to the overall satisfaction score. Kalra et al.13 also described the ODI score as having decreased from the preoperative 56.26 to 11.2 after surgery in 15 patients with rigid tuberculous kyphosis. In a similar study involving 35 patients with fixed sagittal balance, Kim et al.16 reported the improvement of the mean ODI score from 49 to 24 and very good satisfaction in 87% of the patients. Corresponding to the aforementioned results, our study did demonstrate a significant decrease in the mean ODI score after surgery (from 57 to 19) and good functional improvement at the final follow-up evaluation. In addition, when ODI and VAS scores in our study were divided into 2 subgroups based on the etiology of spinal deformity (congenital or acquired), the mean preoperative ODI score was 39.2 in the congeni-
tal cases and 64.7 in the acquired cases, which improved to 24.8 and 16.4, respectively, and the mean preoperative VAS score was 4.2 in the congenital cases and 6.4 in the acquired cases, which improved to 3.3 and 3.2, respectively. This interesting paradoxical finding of greater improvement in the ODI and VAS scores because of higher pain and disability scores preoperatively in the acquired cases may have occurred because in the congenital group most patients presented with deformity, whereas in the acquired group pain was a common presenting feature.

Complications

Bakaloudis et al.\textsuperscript{1} reported 5 medical complications and 1 neurological complication in their series of 12 patients, for a complication rate of 50%. Similarly, Chang et al.\textsuperscript{8} reported complications in 41 of 83 patients in their series, documenting a complication rate of around 50%. Kawahara et al.\textsuperscript{12} reported no neurological complications. However, their study was small, including only 7 patients. Chen et al.\textsuperscript{10} documented complications in 10 of 23 patients. Lenke et al.\textsuperscript{21} reported complications in only 4 of 35 patients in their series of pediatric deformity patients operated on with VCR. Intraoperative neurological deficit occurred in 3 of 23 patients. Wheret et al.\textsuperscript{22} reported complications in 10 of 23 patients. Lenke et al.\textsuperscript{21} reported complications in only 4 of 35 patients in their series of pediatric deformity patients operated on with VCR in pediatric patients was noted in a multicenter trial that included 147 patients.\textsuperscript{20} The authors observed complications in 59% of these patients. Sixty-eight patients had a complication during surgery (the most common being changes in spinal cord monitoring data and blood loss > 2 L), and 43 had a postoperative complication (most often respiratory related). No patient in their series had a complete permanent neurological deficit.

In our series, the overall complication rate was 30.77%. There was a general complication rate of 15%, which included 2 cases of dural tear and 1 case each of implant failure and superficial infection. The neurological complication rate was 7.6%. One patient had a new-onset neurological deficit involving the left L-4 nerve root (quadriceps weakness 3/5), and in 2 other patients the neurological examination results worsened from preoperative Frankel grade C to A in the immediate postoperative period. In these 3 cases, the patients’ deficits all had improved with conservative management at the 3-month follow-up. One patient, however, developed complete paraplegia (congenital kyphoscoliosis, ASO at the L-1 level) that resolved only partially by the final follow-up evaluation. This complication was later found to be attributable to incomplete osteotomy and forced closure with load on the implants, which led to translation within the narrow canal. Neuromonitoring signal changes were detected during manipulation and closure of the osteotomy and reversal measures were taken (as described in detail in Results), but only partial improvement was found in neurological status even at the final follow-up (Frankel grade D). The same patient suffered from screw pullout and construct failure with loss of correction, for which a revision surgery with extension of fixation and redirection of the loosened screw was performed. The learning points from this case were that osteotomy should be completed before closure is attempted to avoid translation over a hinge. In addition, forced closure leads to excessive stress on implants, which can lead to implant failure and pseudarthrosis. Intraoperative neuro-monitoring is an essential tool for a deformity surgeon, as multimodal monitoring acts to warn when damage is done and aids in timely performance of corrective measures. In our case, timely reversal led to partial recovery, which would otherwise have been a complete and irreversible neurological deficit. All other patients in the series had improved with conservative management at the 3-month follow-up.

Limitations

The major limitation of this study is its retrospective design; however, the database for this study was constructed without any hypothesis, and the data were thus collected free from any bias related to the study. Nevertheless, we recognize that prospective data collection would have strengthened the salient points. Another limitation of the study is the small number of cases with varied etiology and a relatively short follow-up. A longer duration of follow-up and larger groups of cases with similar etiology could help elucidate the effect of etiology on correction and morbidity with the technique. Longer follow-up would allow for more observation of the pitfalls of long-segment fusion in the pediatric population, and the effect of residual deformity resulting in mechanical back pain, implant loosening, and loss of correction can be evaluated better. Another limitation of the study was the use of the ODI as an outcome measure, although it has not been validated in the pediatric age group. The ODI was used because there were no validated pediatric outcome measures available in the local language at the time of data collection. Moreover, all of the questions were answered by parents, so a bias of subjective effect on evaluation cannot be ruled out.

Conclusions

In this paper, we described our technique of single-level apical spinal osteotomy (ASO) through a posterior approach for the treatment of severe rigid pediatric kyphoscoliosis. The technique can be performed in small children with good clinical and radiological results and satisfactory correction rates. Although the procedure is technically demanding, it can be reliably executed by any skilled spine surgeon familiar with decompression surgery and instrumentation techniques. It is versatile and effective for correction of rigid kyphoscoliosis in children, the treatment of which to date has proved to be extremely difficult, even in experienced hands.

References

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**Disclosures**

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

**Author Contributions**

Conception and design: Ruparel, Patel, Kundnani. Acquisition of data: Patel, Dusad, Mehta, Kundnani. Analysis and interpretation of data: Patel, Kundnani. Drafting the article: Ruparel, Patel, Kundnani. Critically revising the article: Ruparel, Patel, Kundnani. Reviewed submitted version of manuscript: Ruparel, Patel, Kundnani. Approved the final version of the manuscript on behalf of all authors: Ruparel. Statistical analysis: Kundnani. Administrative/technical/material support: Kundnani. Study supervision: Ruparel, Kundnani.

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