Klippel–Feil Syndrome with Multiple Cervical Anomalies Discovered Following Trauma

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Abstract

The approach to the upper cervical spine is a challenge for surgeons, not only for its complex anatomy but also for the great variety of pathologies. Klippel–Feil syndrome (KFS) is a congenital disease characterized by the fusion of two or more cervical segments and is associated with various musculoskeletal and vascular malformations. However, there is no consensus on the type of surgery, approach, level of fixation or fusion. We report the case of a KFS, associated with multiple anatomical variants, who suffered a traumatic cervical injury and underwent surgical treatment.

Keywords
► Klippel–Feil syndrome
► atlas
► vertebral artery
► cervical fracture
► anterior cervical corpectomy and fusion

Introduction

The management of pathologies of the upper cervical spine is complex, not only due to the anatomy but also due to the diversity of pathologies (craniocervical junction, atlas, axis, ligaments and joints), which generate different approaches for surgical management.¹

Klippel–Feil syndrome (KFS) is a congenital disease characterized by the fusion of two or more cervical segments and is associated with various musculoskeletal and vascular malformations;² up to 50% of patients present the following triad: low posterior hairline, short neck, and decreased range of motion.²-

 Decreased movement due to congenital fusion of the cervical spine, in the long-term, causes stress on the fused segments and produces cervical hypermobility with a consequent increase in the instability of the cervical spine.²

We report the case of a KFS associated with an anomalous course of the vertebral artery, occipitization of C1, luxation of the odontoid process and C3–C4 anterolisthesis, following trauma and surgical management.

Case Report

A 63-year-old male, with a history of a traffic accidents and whiplash syndrome 30 days before admission, presented moderate cervical pain and paresthesias of the left upper limb that lasted 3 days. Due to persistent symptoms and cervical pain (visual analog scale [VAS] 7/10), patient went to the emergency department.

Physical examination revealed cervical pain at the level C2–C6 and paramedial, presence of low-posterior hairline, a wide and short neck with decreased range of motion at flexion, lateral flexion, extension, and rotation movements. Hypoesthesia in dermatomes C3–C5 of the left upper limb, limitation of movement of left shoulder to abduction, C5 myotome muscular strength in left upper limb ⅖ and normal reflexes were found.

Radiological examination showed the cervical spine having lordosis, fusion of C1 and C2 and anterolisthesis C3–C4 (►Fig. 1A, 1C). CT of the cervical spine showed traces of nondisplaced fracture at the level of the right lateral mass of atlas, right lateral luxation of the odontoid process, C3–C4 grade I anterolisthesis and atlanto-occipital assimilation (►Fig. 1D, 1E).

CT angiography of carotid arteries showed a variation in the course of both vertebral arteries. Left vertebral artery was dominant, with a preatlas loop and the entrance to the skull was below C1. The right vertebral artery was hypoplastic, making its entrance to the skull below C1 directly (►Figs. 1F, 1I).

MRI of the cervical spine showed signs of ankylosis between the base of the skull and atlas, atlas and axis; C3–C4 grade I anterolisthesis, C2–C3 protruding hernia with

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posterior osteophytes, and a hypertrophic yellow ligament, which conditioned a compressive myelopathy with edema at this level and spondyloarthrosic changes (►Fig. 2A, 2B).

We decided to perform the anterolateral cervical approach (Smith and Robinson approach). The patient was in supine position with head traction. A left paramedian incision was made and usual dissection was performed. The C3 vertebra was identified and the corpectomy was followed by the placement of the implant. The bony structures were softer and easily resected. Finally, the anterior plate was placed. The identification of the correct levels was the major concern of the procedure.

After surgery, the patient made a gradual recovery with clinical improvement. VAS improved to 3/10 and the postoperative CT scan showed no complications (►Fig. 2C).

Discussion
Management of the upper cervical spine pathology is a challenge due to its complex anatomical characteristics and for the variety of existing surgical classifications and criteria such as those of AO Spine, German Society for Orthopedics and Trauma (DGOU), among others.5–7 Despite these criteria, there is no definitive consensus on the type of surgery, approach, level of fixation or fusion in patients with cervical injury; even less in patients with KFS.8,9

KFS is a genetic disease characterized by fusion of the cervical vertebrae, which generates decreased movement; in the long-term, it produces hypermobility and cervical instability, with a natural history that is not very clear. It presents a wide variety of symptoms and neurological signs, which are intensified when trauma occurs.16 Similar cases like this have been reported; authors like Yusuke, Mishima and others have published cervical fractures associated with KFS.11,12

Our case presented the classic triad: A short neck, decreased range of motion, and low-posterior hairline, which is seen in up to 50% of cases with KFS.8,10,13,14

Patients with KFS also present concomitantly with other congenital anomalies such as spina bifida, Arnold-Chiari, scoliosis, concurrent neuromuscular abnormality, Sprengel deformity, congenital torticollis, cardiovascular abnormalities, abnormalities of the urinary tract, and nervous system abnormalities.2 Regarding vertebral artery anomalies, Patil et al reported an incidence between 3 and 8% of vertebral artery anomalies.15 Few articles have reported abnormalities of the vertebral artery; Ahmad et al reported a case of anomalous vertebral arteries in KFS with occipitalization of the atlas,16 similar to our patient, with the difference that in our case both vertebral arteries entered into the foramen below C1.

There are no exact guidelines indicating the best technique or approach in cervical injuries; however, criteria of instability or presence of neurological deficit in most of the guidelines are taken into account.5–7,9,17,18

When we decided the surgical technique in this patient, we also had to assess the total cost of the materials that would be used (patient did not have insurance). The technique used was the anterolateral cervical approach (Smith and Robinson approach) associated with C3 corpectomy, following the anatomical and surgical steps described in previous reports.19,20

Although there is no clear decision between an anterior and posterior approach, the anterior approach showed better postoperative outcomes, according to neurological recovery, compared with the posterior approach in patients with neurological deficit or cervical myelopathy.20

More studies are needed to determine the best surgical approach in these patients and monitor postoperative outcomes.
Conclusion
The management of cervical spine pathology is complex, as it will depend not only on the type of injury, but also on the comorbidities of each patient. KFS-associated trauma is a challenge for diagnosis and treatment due to the anatomical variations.

Conflict of Interest
None declared.

References
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Fig. 2 (A, B) Preoperative cervical MRI showing the C2–C3 hernia, mild C3–C4 listhesis, and the hypertrophic yellow ligament generating myelopathic changes. (C) Postoperative cervical CT scan showing the final result after anterior cervical corpectomy and fusion (ACCF).