Embryological considerations and evaluation of congenital anomalies of craniovertebral junction: A single-center experience

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ABSTRACT

Objectives: Craniovertebral junction (CVJ) abnormalities constitute a group of treatable neurological disorders, especially in the Indian subcontinent. Thus, it is essential that clinicians should be able to make a precise diagnosis of abnormalities and rule out important mimickers on multidetector computed tomography (MDCT) as this information ultimately helps determine the management, prognosis, and quality of life of patients. CVJ is the most complex part of the cervical region. Congenital malformations of this region can cause serious neurological deficit and require a surgical intervention. The present study was undertaken to know the embryological basis of the CVJ and to identify commonly observed congenital CVJ abnormalities, their frequency, and mode of presentation.

Materials and Methods: Diagnosed cases of CVJ anomalies on dynamic MDCT head were reviewed at a tertiary care center between January 2014 to December 2019. Type of anomaly, clinical presentation, and associated malformations were recorded. Different types of variations were expressed in terms of percentage. Results: Congenital anomalies were seen in 42 cases. Fifteen types of anomalies were detected. Anomalies were either singly or in combination. The CVJ anomalies were more common in young adults (28%), almost equal in both sexes. The most common anomaly was basilar invagination (52.3%), followed by atlanto-occipital assimilation (33.3%), and Arnold–Chiari malformation is the most common soft tissue anomaly. In fourteen cases, additional anomalies of other vertebrae were present. The most common symptoms were weakness of extremities, neck pain, paresthesia, torticollis, and gait disturbances. About 28 patients got improved, 8 patients had residual deficit as that of preoperative status, and 4 patients got deteriorated after surgery. Conclusion: Congenital CVJ anomalies, though rare, are fatal. CVJ abnormalities constitute an important group of treatable neurological disorders with diagnostic dilemma. The atlantodental interval is the most important preoperative prognostic marker. Dynamic CT imaging can provide additional useful information to the diagnosis of CVJ instability. To prevent long-term neurological problems, early diagnosis and treatment of congenital bony CVJ anomalies is important.

Keywords: Basilar invagination, Cranio metric parameters, Craniovertebral junction anomalies, Multidetector computed tomography, Sclerotomes

INTRODUCTION

Craniovertebral junction (CVJ) consists of basilar part of occipital bone, atlas, and axis vertebra. Being the transit zone between cranium and spine, it is the most complex and dynamic area of the cervical region. It has complex bony anatomy and is related with major neurovascular structures. It shows extensive variability in morphology also. Congenital malformations associated with this region have a potential of causing serious neurological and vascular deficit and

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may require surgical intervention. The incidence of different types of CVJ anomalies varies with demographic regions and genetic factors. CVJ anomalies are more frequently found in the Indian subcontinent than anywhere else in the world [1]. These osseous anomalies can manifest with abnormal cerebrospinal fluid dynamics [2]. Multidetector computed tomography (MDCT) is the most commonly used modality to assess the CVJ [3]. To identify different anomalies, it is important to understand the embryology and developmental anatomy of the region. It is important to rule out mimickers on MDCT as this information ultimately helps determine the management, prognosis, and quality of life of patients with CVJ anomalies. Hence, the current study was undertaken to know the embryological basis of the CVJ and to study commonly observed congenital CVJ abnormalities, their frequency, and mode of presentation.

Epidemiology of craniovertebral junction anomalies

The incidence of different types of CVJ anomalies varies with demographic environment and ill-defined genetic factors. CVJ anomalies are more frequently found in the Indian subcontinent than anywhere else in the world. They are common in all age groups and almost equal in both sex groups. The clinical features are often delayed up to 2nd or 3rd decades, since they are subtle and often missed. In the current study, we had screened 78 cases with suspected anomalies of CVJ based on clinical features and found 42 congenital anomalies of CVJ, which was rare and fatal.

Materials and methods

A retrospective, cross-sectional study was conducted at a tertiary care center based on analysis of CT head reports between January 2014 to December 2019. Diagnosed cases of CVJ anomalies on CT head were reviewed. Total 42 patients with bony congenital malformations of age 8 years and above were included in the study. They were divided into six groups according to the age in decade. Side and type of anomaly, clinical presentation, and associated malformations were recorded. Normal CT reports, CT of patients with a history of trauma, tumors, tuberculosis, and rheumatoid arthritis were excluded. Dynamic craniometric measurements on MDCT included Chamberlain’s line, McRae’s line, McGregor’s line, Wackenheim’s clivus canal line, and Welcher’s basal angle. Magnetic resonance imaging (MRI) was done in suspected cases of neurological involvement. After complete clinical examination, the sensory level, the motor level, and the reflex level were found and the diagnosis of CVJ anomalies was attained. The other associated clinical features such as head tilt, short neck, webbed neck, downbeat nystagmus, cranial nerve deficits, and features of congenital syndromes such as Down syndrome, Morquio syndrome, and Klippel–Feil syndrome were taken into account. Atlantodental interval is the single important and reliable marker to assess the prognostic outcome. Various bony anomalies such as defects in the arches of atlas, assimilation of atlas, block vertebrae, basilar invagination (BI), atlanto-occipital dislocation, and odontoid fracture were studied. Among the congenital bony CVJ anomalies, congenital atlanto-axial subluxation, BI, platybasia, occipitalization of atlas, and defect in the arches of atlas were studied. Among the congenital soft tissue CVJ anomalies, Arnold–Chiari malformations and syringomyelia were studied. All the reducible CV junction anomalies are better treated with surgical modality. For surgical repair under fluoroscopy guidance, lateral mass screw on C1 and pars or pedicle screw on C2 were placed bilaterally. Occipital plate was used for occipitocervical fusion. The irreducible anomalies are immobilized with Philadelphia collar or halo brace and managed conservatively. Different types of variations were expressed in terms of percentage. The study was conducted in accordance with the Declaration of Helsinki and was approved by the Institutional Ethics Committee (IEC), and informed consent was obtained from all patients before their enrollment in this study (IEC, St. John’s Hospital; IEC Approval Reference Number: SJH/7/2014; IEC Approval Date: January 12, 2014).

Statistical analysis

The findings were tabulated using Microsoft Excel 2010 and statistical analyses were conducted using SPSS Statistical Package (version 20.0), IBM SPSS Statistics for Windows, V.20.0, IBM Corp., Armonk, New York, USA.

Results

Congenital CVJ anomalies were seen in 42 cases. Fifteen types of anomalies were detected [Table 1]. Anomalies seen were either singly or in combination [Figures 1 and 2]. The most common anomaly was BI seen in 52.3% of cases; BI was seen in combination with atlanto-occipital assimilation, occipital condylar hypoplasia, ponticulus posticus, and atlantoaxial assimilation [Table 2]. In fourteen cases, associated anomalies of other vertebrae were present [Table 2]. Maximum of 12 cases were detected in the age group of 11–20 years. The most common symptoms were weakness of extremities, neck pain, paresthesia, torticollis, and gait disturbances.

In the current study, age distribution was from 8 to 60 years and most of the patients were diagnosed in the

| Table 1: Incidence of craniovertebral junction anomalies in the current study |
|---|---|
| Type of CVJ anomaly | n (%) |
| BI | 22 |
| AOA | 14 |
| Incomplete posterior arch of atlas | 6 |
| C2-C3 fusion | 4 |
| Proatlas/condylus tertius | 4 |
| Os terminale | 4 |
| OCH | 2 |
| PP | 1 |
| Hypertrophy of anterior arch of atlas | 2 |
| CAA | 2 |
| Platybasia | 2 |
| Hypoplasia of posterior arch of atlas | 2 |
| OS odontoideum | 2 |
| Absent lateral mass of atlas | 1 |
| C1-C2 dislocation | 1 |

CVJ: Craniovertebral junction, BI: Basilar invagination, AOA: Atlanto-occipital assimilation, OCH: Occipital condyle hypoplasia, PP: Ponticulus posticus, CAA: Complete atlantoaxial assimilation
second decade with male predominance. Patients in the current study presented with complaints such as weakness of extremities (52%), neck pain (32%), paresthesia (8%), torticollis (4%), and gait disturbances (4%). In children with unilateral atlas assimilation, torticollis was a presenting symptom.

Skeletal deformities were found in 4 patients (3 had kyphosis and 1 had scoliosis), spastic quadriapresis in 4 patients, sensory disturbances involving the anterolateral spinocerebellar tract and posterior column in 38 patients, autonomic disturbances in 28 patients (isolated bladder involvement in 18 patients, combined bladder and bowel involvement in
The vertebral column develops in six separate but overlapping phases. The first phase is gastrulation and the formation of the mesoderm and notochord. In the second phase, somites are formed. During the third phase, somites are reorganized to form dermomyotome and sclerotome. There is resegmentation of the somites to form the definitive vertebrae in the fourth phase. This is followed by vertebral chondrification in the fifth and vertebral ossification in the sixth phase [4]. There are about 42 pairs of somites in the 4th week of development [5]. There are 4 occipital somites, 8 cervical, 12 thoracic, 5 lumbar, 5 sacral, and 8–10 coccygeal pairs of somites. The cells of sclerotome migrate ventromedially around notochord to form primitive vertebral bodies [6]. The first and second occipital sclerotomes form the basilar part of occipital bone. Caudal part of the 4th occipital somite fuses with cranial half of the 5th somite or first cervical to form proatlas sclerotome. The CVJ abnormalities are caused by abnormalities of resegmentation of the proatlas. The proatlas forms the anterior tubercle of the clivus, tip of the dens, and the apical ligament [7]. The neural arch of the proatlas forms the anterior margin of the foramen magnum, occipital condyle, the lateral atlantal masses, and the superior portion of the posterior arch of the atlas. The axis body is formed by the centrum and the neural arch forms the facets and the posterior arch of the axis vertebra [8]. The tip of the odontoid process is derived from the proatlas [9]. A separate ossification center appears for the tip of the odontoid at 3 years of age and fuses with the rest of the dens by 12 years of age. Resegmentation of sclerotome is under the control of regulatory genes such as Hox and Pax [10]. Both MTHFR 677C→T polymorphism and higher T allele frequency have significant associations with AAD. Genotypes with single base substitutions, deletions, or duplication of part of the COL2A1 gene have been advocated in CVJ anomalies. Cervical spine radiographs in these patients have demonstrated apparent atlantoaxial instability in correlation with odontoid hypoplasia or Os odontoideum [11]. Pax genes are involved in vertebral formation and contribute to the development of the early nervous system [12].

Clinical presentation

The most interesting feature is the diversity of clinical feature as a result of compromise of lower brainstem, cervical spinal cord, cranial nerves, cervical roots, and vascular supply. The congenital anomalies are associated with abnormal physical appearance such as head tilt, short neck, low hairline, and limited neck movements (Klippel–Feil syndrome). The most frequent symptom is the suboccipital neck pain, radiating to the cranium. The features of myelopathy may present in the form of monoparesis, hemiparesis, paraparesis, and ascending pattern of quadriparesis. Central cord syndrome is seen in children with BI. Sensory abnormalities are manifested as neurological deficits related to the anterolateral and posterior column dysfunction. Brainstem and cranial nerve deficits can cause abnormalities such as dysphagia, nasal regurgitation, change in voice, loss of facial sensation, and sleep apnea. Downbeat nystagmus is a cardinal sign, more characteristically seen with strictly compressive CVJ lesions. The excessive mobility of an unstable CVJ can cause trauma to the anterior spinal artery and the perforating vessels of the upper cervical cord and medulla oblongata producing features of verteobasilar insufficiency such as vertigo, syncopal attacks, and transient loss of consciousness.

Congenital CVJ al anomaly is a rare condition all over the world and most studies have reported few cases seen over a long period [13]. The current study demonstrated anomalies in 42 patients over a period of 6 years. A study reported 189 patients with the same anomalies seen in 10 years [14]. In another study, 23 cases were reported from a hospital in northern India in 5 years [15]. The most common congenital anomaly observed in the present study was BI in 22 cases. The congenital anomalies are associated with abnormal physiology of the CVJ. The congenital anomalies are associated with abnormal physiology of the CVJ.

Distribution of surgical outcome

The neurological outcome was assessed in these 42 patients by thorough clinical examination of motor power under MRC Grade. About 28 patients got improved, 8 patients had residual deficit as that of preoperative status, and 4 patients got deteriorated after surgery at 1-month follow-up. About 34 patients had improved, 5 remained static, and 3 patients got worsened at the end of 3 months of follow-up. About 37 patients had improved, 4 patients remained static, and patients got deteriorated at 6 months of follow-up.

The CVJ anomalies are more common in young adults (28%), almost equal in both sexes. BI (52.3%) is the most common congenital bony anomaly, and Arnold–Chiari malformation is the most common soft tissue anomaly. The patients with increased atlantodens interval 3–5 mm showed 77% improvement after surgery.

DiSCUSSION

Embryology of craniovertebral junction

The vertebral column develops in six separate but overlapping phases. The first phase is gastrulation and the formation of the mesoderm and notochord. In the second phase, somites are formed. During the third phase, somites are reorganized to form dermomyotome and sclerotome. There is resegmentation of the somites to form the definitive vertebrae in the fourth phase. This is followed by vertebral chondrification in the fifth and vertebral ossification in the sixth phase [4]. There are about 42 pairs of somites in the 4th week of development [5]. There are 4 occipital somites, 8 cervical, 12 thoracic, 5 lumbar, 5 sacral, and 8–10 coccygeal pairs of somites. The cells of sclerotome migrate ventromedially around notochord to form primitive vertebral bodies [6]. The first and second occipital sclerotomes form the basilar part of occipital bone. Caudal part of the 4th occipital somite fuses with cranial half of the 5th somite or first cervical to form proatlas sclerotome. The CVJ anomalies are caused by abnormalities of resegmentation of the proatlas. The proatlas forms the anterior tubercle of the clivus, tip of the dens, and the apical ligament [7]. The neural arch of the proatlas forms the anterior margin of the foramen magnum, occipital condyle, the lateral atlantal masses, and the superior portion of the posterior arch of the atlas. The axis body is formed by the centrum and the neural arch forms the facets and the posterior arch of the axis vertebra [8]. The tip of the odontoid process is derived from the proatlas [9]. A separate ossification center appears for the tip of the odontoid at 3 years of age and fuses with the rest of the dens by 12 years of age. Resegmentation of sclerotome is under the control of regulatory genes such as Hox and Pax [10]. Both MTHFR 677C→T polymorphism and higher T allele frequency have significant associations with AAD. Genotypes with single base substitutions, deletions, or duplication of part of the COL2A1 gene have been advocated in CVJ anomalies. Cervical spine radiographs in these patients have demonstrated apparent atlantoaxial instability in correlation with odontoid hypoplasia or Os odontoideum [11]. Pax genes are involved in vertebral formation and contribute to the development of the early nervous system [12].

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assimilation. Another study also reported the similar combination in 14.5% of cases [17]. The second most common anomaly observed in the current study was atlanto-occipital assimilation in 14 cases (33.3%). Mwang’ombe and Kirongo reported occipitalization of the atlas (28%) [18]. Congenital fusion of the atlas with the occiput is one of the most common anomalies of the CVJ, with a prevalence rate from 0.08%–2.8% in the general population [19]. Sometimes, two or more vertebrae may fuse to giving rise to Klippel–Feil syndrome and occipitalization of atlas [20]. The current study recorded fusion in fourteen cases. Other atlas anomalies included two cases each of hypertrophied anterior arch, hypoplastic posterior arch, atlantoaxial fusion, six cases of incomplete posterior arch, absence of lateral mass in one case, and ponticulus posticus in one case. Defects in posterior arch of atlas are ten times more common than defects of anterior arch. Incomplete posterior arches are commonly seen. The defect never closes with age [21]. Platybasia was noted in two cases. Hypoplasia of occipital condyles was seen in two cases in the present study. Hypoplastic condyles results in flattening of skull base which may lead to BI. According to Menezes and VanGilder, condylus tertius or median occipital condyle is observed due to failure of proatlas integration [22]. Os odontoideum is frequently associated with condylus tertius [23]. In the current study, condylus tertius or proatlas was observed in four cases. Os odontoideum was noted in two cases. In this, odontoid process may remain separate from the body of the axis partially or completely [24]. In four cases, Os terminale was seen.

In a study of 70 patients by Menezes, 85%–90% age of presentation was between the first and second decade of life [25]. Other studies reported the same age distribution with male predominance and third decade presentation [17]. Menezes observed spastic quadripareisis as a presenting symptom in 80% of the patients and lower cranial nerve palsies in 33% of the patients [26]. Other studies reported progressive weakness of the extremities due to syringomyelia as the most common mode of presentation in atlantoaxial instability [27]. The various surgical treatment options for CVJ anomalies are described as follows: (1) foramen magnum decompression, (2) suboccipital craniectomy and removal of posterior arch of atlas, (3) C1–C2 wiring done by sublaminar titanium wires, (4) C1–C2 fixation done by C1 lateral mass and C2 pedicle screw fixation, (5) occipitocervical fusion is done by stainless steel or titanium contoured rod or loop with screw fixation, and (6) transoral odontoidectomy is done by transpalatal, transpharyngeal routes to relieve the ventral compression.

**CONCLUSION**

Congenital CVJ anomalies, though rare, are fatal. CVJ abnormalities constitute an important group of treatable neurological disorders with diagnostic dilemma. Craniometrical parameters, as well as the visualization of the facets location, may change significantly according to the neck position. The atlantodental interval is the most important preoperative prognostic marker. Dynamic CT can provide additional useful information to the diagnosis of CVJ instability. Early diagnosis and treatment form the mainstay in preventing long-term neurological complications. MDCT is the investigation of choice for diagnosis and planning the management. CT complemented with MRI is recommended for associated neurological involvement.

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**Conflicts of interest**

There are no conflicts of interest.

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